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TRANSACTIONS

Volume XX

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THE STROUD FORAY

May 11th-15th, 1934

By E. M. WAKEFIELD

THE Spring Foray for 1934 was held during the week-end May 11th-15th, with headquarters at the Amberley Ridge Hotel, near Stroud.

The first day's excursion was to Cranham Woods, between Painswick and Cheltenham. A start was made from the hotel at 10.15 a.m., and the party travelled by bus as far as the King William Hotel, near Cranham Corner. Some little time was spent in the woods behind the hotel in a search for *Pyrola*, which had been found there during the Painswick Foray in 1920, attacked by its rust, *Chrysomyxa Pyrolae*. Unfortunately, however, the plant appears to have now become almost extinct in that locality, owing to the use of the wood by picnic parties.

A move was then made towards Cranham Corner, and the woods between there and Cranham Village were explored. Owing to the dry winter and spring the larger fungi were very scarce. The most noteworthy find was Sarcosphaera coronaria, which occurred in abundance in certain spots, growing amongst moss on the ground and at the foot of banks beside the paths. A few rusts and other microfungi were

secured, but nothing of any special interest.

On the following day, Sunday, Woodchester Park was the objective. Sufficient private cars were available, and by a steep and narrow rutty lane the party arrived in procession at the gates of the Park, and were admitted by the lodge-keeper. The Park was found to be a place to rejoice a botanist's heart, and many remarked how good it would be for fungi in autumn. At this time, like the rest of the country, it had suffered from the drought, but at the same time the variety of vegetation provided many more fungi than had been found on the previous day. Some cherries were found badly attacked by leaf-curl (Taphrina minor). Among other microfungi collected were Entyloma Ranunculi, abundant on Ranunculus Ficaria, and Endophyllum Euphorbiae-sylvaticae on Euphorbia amygdaloides. Basidiomycetes were also more in evidence, owing partly to moisture provided near a chain of lakes, and partly to the presence of plenty of rotting wood. Eichleriella spinulosa, found during the Painswick Foray, was again collected, while of the larger fungi Tricholoma gambosum, Coprinus ephemerus, and Polyporus dryadeus may be noted.

Monday was devoted to the exploration of Cirencester Park. Here again, however, fungi were very few. A good group of Morchella rotunda was found, fine large specimens, some of which were taken

for a culinary test. There was little of any special interest, but *Penio-phora longispora* and *Aleuria umbrina* were among the less common

species recorded.

During the week-end a meeting was held to consider proposals for the 1935 Spring Foray. After considerable discussion Derbyshire (Buxton or Matlock) was proposed, and Tunbridge Wells was suggested as an alternative should it not be possible to arrange for a Derbyshire meeting. The final decision was left with the Council.

Miss L. Hunter, of Toronto, gave an informal account of her work on the genus *Milesia*. Mr Petch gave an interesting talk on some entomogenous fungi found in caves, and Dr Alex. Smith showed specimens of the very rare *Puccinia Bulbocastani* on *Carum Bulbocastanum*. At the close of the meeting votes of thanks were accorded, and special thanks were expressed to Mr E. M. Day, who had made all arrangements for the very pleasant meeting. Members regretted that owing to illness Mr Day was unable to join in the excursions.

The Secretary is indebted especially to Mr Ramsbottom, Mr Pearson, Dr Alex. Smith, Mr T. Petch, and Mr E. W. Mason for

assistance in compiling the subjoined list.

Complete List of Species found during the Foray

P. = Cranham Woods, near Painswick; C. = Cirencester Park; W. = Woodchester Park; A. = Amberley Hotel grounds.

HYMENOMYCETES

Armillaria mellea (Vahl) Fr., rhizomorphs, C. Tricholoma gambosum Fr., W. Mycena amicta Fr., P. Marasmius globularis Fr., C., dryophilus (Bull.) Karst., C. Panus stipticus (Bull.) Fr., W. Schizophyllum commune Fr., W. Nolanea pascua (Pers.) Fr., W. Pholiota praecox (Pers.) Fr., W., C., mutabilis (Schaeff.) Fr., W. Hebeloma crustuliniforme (Bull.) Fr., C. Galera tenera (Schaeff.) Fr., C., P. Naucoria melinoides Fr., W. Tubaria furfuracea (Pers.) W. G. Sm., P. Crepidotus mollis (Schaeff.) Fr., W. Psalliota campestris (Linn.) Fr., C. Hypholoma fasciculare (Huds.) Fr., P., W., C., velutinum (Pers.) Fr., P., appendiculatum (Bull.) Fr., P., W. Psilocybe foenisecii (Pers.) Fr., W. Psathyrella disseminata (Pers.) Fr., P. Panaeolus campanulatus (Linn.) Fr., W. Coprinus micaceus (Bull.) Fr., C., P., velox Godey, C., ephemerus (Bull.) Fr., Polyporus squamosus (Huds.) Fr., W., C., sulphureus (Bull.) Fr., C., dryadeus (Pers.) Fr., W., radiatus (Sow.) Fr., W., fumosus (Pers.) Fr., P., adustus (Willd.) Fr., W., C., caesius (Schrad.) Fr., W., P. Fomes ulmarius (Sow.) Fr., W., C., ferruginosus (Schrad.) Mass., W.

Ganoderma applanatum (Fr.) Pat., W.
Polystictus versicolor (Linn.) Fr., P., W., C., abietinus (Dicks.) Fr., W.

Polystictus versicolor (Linn.) Fr., P., W., C., abietinus (Dicks.) Fr., W.
Irpex obliquus (Schrad.) Fr., W.
Lenzites betulina (Linn.) Fr., W.
Trametes gibbosa (Pers.) Fr., W., mollis (Sommerf.) Fr., W.
Daedalea quercina (Linn.) Fr., W.
Stereum spadiceum Fr., W., rugosum (Pers.) Fr., W., hirsutum (Willd.) Fr., W., purpureum Pers., W., C.
Corticium laeve Pers., W., C., lividum (Pers.) Fr., C., porosum B. & C., C., praetermissum (Karst.) Bres., W., lactescens Berk., W.
Peniophora longispora (Pat.) v. H. & L., C., cremea Bres., W., C., velutina (DC.)
Cooke, W., hydnoides Cooke & Mass., W., C., cinerea (Fr.) Cooke, W., C.
Auricularia auricula-Judae (Linn.) Schroet., W.
Tremella mesenterica (Retz.) Fr., W.

Tremella mesenterica (Retz.) Fr., W. Exidea nucleata (Schw.) Rea, P., Thuretiana (Lév.) Fr., C. Eichleriella spinulosa (B. & C.) Burt., W.

GASTEROMYCETES

Lycoperdon perlatum Pers., C.

UREDINALES

Uromyces Ficariae (Schum.) Lév., W., Acetosae Schroet., W., Alchemillae (Pers.) Lév., C., Scillarum (Grev.) Wint., P., C., Dactylidis Otth, aecidium on Ranunculus, P.

Puccinia Anemones Pers., P., C., Violae (Schum.) DC., P., pulverulenta Grev., W., Saniculae Grev., P., Celakovskiana Bubák, W., obtegens (Link) Tul., W., Hieracii (Schum.) Mart., P., Taraxaci Plowr., P., Buxi DC., C.

Kuehneola albida (Kuehn.) Magn., W., C. Ochropsora Sorbi (Oud.) Diet., aecidium on Anemone, P.

Endophyllum Euphorbiae-sylvaticae Wint., on Euphorbia amygdaloides, P., W.

USTILAGINALES

Entyloma Ranunculi (Bon.) Schroet., on Ranunculus Ficaria, W. Urocystis Anemones (Pers.) Schroet., P., C.

PYRENOMYCETES

Podosphaera Oxyacanthae (DC.) de Bary, P. Sphaerotheca pagnosa (Wallr.) Lév., P. Microsphaera Berberidis (DC.) Lév., on Mahonia Aquifolium, A.

Erysiphe graminis DC., W.
Nectria Peziza (Tode) Fr., C., sanguinea (Bolt.) Fr., W.
Hypomyces rosellus (A. & S.) Tul., W., aurantius (Pers.) Tul., C.

Leptospora ovina (Pers.) Fuck., W.

Stigmatea Robertiana Fr., P.

Eutypa Acharii Tul., on Acer, W., lata (Pers.) Tul., C., spinosa (Pers.) Tul., C.

Botryosphaeria Dothidea (Moag.) Ces. & de Not., on Rosa arvensis, P.

Diatrypella quercina (Pers.) Nits., W.
Diatrypella quercina (Hoffm.) Fr., on Fagus, W.
Ustulina vulgaris Tul., P., W., C.
Hypoxylon coccineum Bull., W., fuscum (Pers.) Fr., on Corylus, W., C.
Daldinia concentrica (Bolt.) Ces. & de Not., W.

Xylaria Hypoxylon (Linn.) Grev., C.

HYSTERIALES

Dichaena faginea Fr., C.

DISCOMYCETES

Morchella rotunda (Pers.) Boud., C. Aleuria umbrina Boud., C. Sarcosphaera coronaria (Jacq.) Boud., P. Cheilymenia coprinaria (Cooke) Boud., P. Ascobolus stercorarius (Bull.) Schroet., P. Dasyobolus immersus (Pers.) Sacc., P. Pyronema omphalodes (Bull.) Fuck., C. Taphrina minor Sadeb., on Cerasus, W. Calycella citrina (Hedw.) Quél., W. Bulgaria inquinans (Pers.) Fr., C. Dasyscypha virginea (Pers.) Fr., P. Rhytisma acerinum (Pers.) Fr., P.

PHYCOMYCETES

Cystopus candidus (Pers.) de Bary, on Arabis, A. Plasmopara pygmaca Schroet., on Anemone, W., C. Peronospora alsinearum Casp., W., calotheca de Bary, C. Empusa Muscae Cohn, A.

DEUTEROMYCETES

Stagonospora Curtisii (Berk.) Sacc., A.
Septoria Rubi West., P.
Ceuthospora phacidioides Grev., on Ilex, C.
Gloeosporium Helicis (Desm.) Oud., P.
Leptostroma filicinum Fr., W.
Gliocladium penicillioides Corda, on Didymium squamulosum, W.
Botrytis cinerea Pers., W.
Cylindrodendrum album Bon., on Alder catkins, W.
Ovularia obliqua (Cooke) Oud., W.
Beauveria Bassiana (Bals.) Vuill., on weevils, P.
Ramularia acris Lindr., on Ranunculus repens, W., Calthae Lindr., W.
Bispora monilioides Corda, P., C.
Tilachlidium tomentosum (Schrad.) Lindau, on Trichia varia, C.
Isaria farinosa (Holmsk.) Fr., on pupae, P.

MYCETOZOA

By T. PETCH

Ceratiomyxa fruticulosa Macbr., W.
Physarum nutans Pers., C., viride Pers., C.
Fuligo septica Gmel., W.
Didymium squamulosum Fr., W.
Stemonitis fusca Roth, W.
Comatricha typhoides Rost., W., C.
Lycogala epidendrum Fr., P., W., C.
Trichia affinis de Bary, C., persimilis Karst., C., varia Pers., C., decipiens Macbr., W., C.
Arcyria denudata Wettst., P., W., C., incarnata Pers., W.



THE NORWICH FORAY

October 1st-6th, 1934

By E. M. WAKEFIELD

The thirty-eighth Autumn Foray and Annual General Meeting was held at Norwich during the week October 1st-6th, at the invitation of the Norfolk and Norwich Naturalists' Society. By kind permission of the Castle Museum Committee, headquarters were at the Castle Museum, where a room for meetings was available, and space for the exhibition of specimens. Here a large number of members and visitors assembled on the Monday evening, and were met by members of the Norfolk and Norwich Naturalists' Society. Miss Geldart, Vice-President of the Norfolk and Norwich Naturalists, made a short speech of welcome, to which Mr Ramsbottom replied on behalf of

the British Mycological Society.

On the morning of October 2nd an early start was made by special coach in the direction of Cromer. Stopping at Northrepps Hall, the party were welcomed by Mrs and Miss Gurney, and after a glance through the gardens proceeded to work through Overstrand Woods, mixed woods with a considerable number of conifers (spruce and silver fir). Hymenochaete Mougeotii was soon discovered. It was first found in these woods some years ago. One of the most striking finds was a fallen trunk of silver fir which was completely covered with Polyporus benzoinus. Seen thus in a fresh and actively growing state the fungus is very different in appearance from the dark-coloured, shrivelled, dried specimens. In the growing state it has a conspicuous, swollen, white margin, immediately behind which there is a bright brown zone. Some of the firs in these woods were obviously failing, and as both Armillaria mellea and Fomes annosus were commonly present it is probable that these two fungi were doing considerable damage. After traversing these woods the party was picked up again by the coach, and taken on to woods at West Runton, where Hymenochaete Mougeotii was again turned up. Rain began soon after lunch and became very heavy towards tea time, so that after tea the party was glad to get into the coach and return to Norwich.

Wednesday's excursion was to Westwick and North Walsham, starting at 11.0 a.m. Westwick proved very rich in fungi, and most members felt they could have profitably spent the day in the area where they were first put down. The long subsequent walk to North Walsham yielded very little, but tea at the end, kindly provided by Mr and Mrs J. B. Brookes, was most welcome. A brief roadside survey before actually entering the woods at Westwick yielded

numerous Agarics and Boleti, including Boletus viscidus, B. elegans, Paxillus atrotomentosus, and other conifer-loving species. In the woods Sparassis crispa was found, and other noteworthy records were Russula claroflava, R. sphagnophila, R. paludosa Britz., and Puccinia Hydrocotyles. The woods at North Walsham yielded Cordyceps capitata and C.

ophioglossoides.

Thursday was spent chiefly in woods and heathy ground at Stratton Strawless. Almost at once a quantity of the curious, sterile mycelium known as Anthina flammea was found amongst dead leaves. By the roadside Pulvinula constellatio was secured, and in the woods the most noteworthy finds were Sparassis crispa, Cyathus striatus, Helvella lacunosa, Russula sphagnophila, Cortinarius phoeniceus, Volvaria bombycina, Leptonia sarcita, and Polyporus brumalis. After tea at Cawston a short visit was paid to Buxton Heath, the only known locality for Bovistella paludosa. One old specimen was secured, but the visit was evidently too late for getting this fungus in good condition. On the Heath were found also Entoloma Bloxamii, and Leptonia formosa.

On Friday, October 5th, the party spent the morning in Sprowston and Plumstead Woods, and the afternoon at Framingham Chase and Framingham Manor Woods. At Framingham Puccinia Antirrhini was noticed in great quantity in a lodge garden, and at the same place was found a rather meagre specimen of Cronartium asclepiadeum Fr., on Tropaeolum majus. A small party made a special expedition to Wheatfen Broad, Surlingham, and added a number of records to the

list.

The Annual General Meeting was held on the Tuesday evening, when the Officers and Council for the ensuing year were elected. Dr Malcolm Wilson was unanimously elected as President for 1935. Dr B. Barnes and Mr F. G. Gould are Vice-Presidents. The Treasurer, Secretaries and Editors remain as before. New members of Council, to replace those retiring under the Rules, are Mr C. G. C. Chesters, Mr H. J. Howard, and Mr H. H. Knight.

The list of members of the Plant Pathology Committee for 1935, elected by the Committee, was read and confirmed. New members

are Mr Cartwright, Mr Ogilvie, and Miss K. Sampson.

A very warm invitation had been received from representative Irish botanists to hold the 1935 Autumn Foray at Killarney. It was felt, however, that southern England was due for a visit, and, moreover, the visit to Belfast had been comparatively recent. For this reason Dr O'Connor, who had transmitted the invitation, was asked if it might be held over for a future date. For 1935 it was decided to try to arrange a Foray in Devon.

The Presidential Address, on "Induced Variation", was delivered by Dr Barnes on Wednesday evening, October 3rd. On Thursday evening Mr H. Ramage (visitor) gave a talk on the Mineral Content of



Fungi, illustrated by lantern slides showing spectra. Miss Cayley gave a preliminary account of some work she had been doing with mushrooms, especially the species Psalliota campestris, hortensis, and arvensis.

On Friday evening Mr Rea contributed one of his informal and instructive talks on the more interesting species of fungi which had

been found during the week.

The meeting ended with hearty votes of thanks to all the landowners concerned, to the Norfolk and Norwich Naturalists, and to the Committee and Staff of the Castle Museum, and particularly to Mr G. J. Cooke, who had made all the local arrangements and who with Mrs Cooke had contributed so much enthusiasm and energy to the organisation of a most enjoyable foray.

For assistance in compiling the attached list of species the Secretary is indebted to all members present, but especially to Mr Rea, Mr Ramsbottom, Mr Pearson, Mr Petch, Mr E. W. Mason, and Dr

Alex. Smith.

Complete List of Species found during the Foray

 \mathcal{N} . = Northrepps Hall and Overstrand Woods; R. = West Runton; W. = Westwick and North Walsham; S. = Stratton Strawless; B. = Buxton Heath; P. = Sprowston and Plumstead Woods; F. = Framingham Chase; C. = Cawston; G. = Wheatfen Broad, Surlingham.

HYMENOMYCETES

Amanita verna (Lam.) Fr., N., R., W., phalloides (Vaill.) Fr., W., P., S., porphyria (A. & S.) Fr., S., mappa (Batsch) Fr., R., W., S., P., and var. alba (Gill.) Rea, S., muscaria (Linn.) Fr., R., W., S., and var. formosa Fr., S., spissa Fr., W., S., rubescens (Pers.) Fr., N., R., W., S., P., G.
Amanitopsis fulva (Schaeff.) W. G. Sm., W., P., S., G.
Lepiota procera (Scop.) Fr., R., S., rhacodes (Vitt.) Fr., S., echinella Quél. & Bern., P., G., felina (Pers.) Fr., N., R., cristata (A. & S.) Fr., W., S., granulosa (Ratsch) Fr. M. W. amianthina (Scop.) Fr. W.

(Batsch) Fr., N., W., amianthina (Scop.) Fr., W.

Armillaria mellea (Vahl) Fr., N., W., P., S., G., mucida (Schrad.) Fr., on beech

and oak, S.

Tricholoma resplendens Fr., F., fulvum (DC.) Fr., S., albobrunneum (Pers.) Fr., W., S., rutilans (Schaeff.) Fr., N., R., W., P., acerbum (Bull.) Fr., W., nudum (Bull.) Fr., N., cinerascens (Bull. non Fr.) Quél., S., P., grammopodium

(Bull.) Fr., S., melaleucum (Pers.) Fr., W., S., sordidum (Schum.) Fr., W.
Clitocybe nebularis (Batsch) Fr., S., clavipes (Pers.) Fr., W., S., aurantiaca (Wulf.)
Studer, N., R., W., F., G., and var. albida (Gill.) Rea, W., B., odora (Bull.)
Fr., W., rivulosa (Pers.) Fr., W., candicans (Pers.) Fr., R., infundibuliformis
(Schaeff.) Fr., N., R., W., G., inversa (Scop.) Fr., P., flaccida (Sow.) Fr., W.,
G., suaveolens (Schum.) Fr., N., W., F., P., ditopus Fr., W., vibecina Fr., N., S.

Laccaria laccata (Scop.) B. & Br., N., W., S., P., G., and var. amethystina (Vaill.) B. & Br., N., S., F.

Collybia radicata (Relh.) Berk., W., S., F., P., platyphylla (Pers.) Fr., R., N., W., maculata (A. & S.) Fr., N., R., W., P., G., distorta Fr., W., butyracea (Bull.) Fr., W., cirrhata (Schum.) Fr., N., S., tuberosa (Bull.) Fr., R., W., S., F., atrata Fr., S., protracta Fr., N.

Mycena pelianthina Fr., \mathcal{N} ., rubro-marginata Fr., R., P., pura (Pers.) Fr., \mathcal{N} ., W., polygramma (Bull.) Fr., S., galericulata (Scop.) Fr., N., R., S., P., G., polygramma (Bull.) Fr., S., alcalina Fr., N., ammoniaca Fr., N., R., S., P., G., vitilis Fr., R., amicta Fr., F., Iris Berk., P., F., sanguinolenta (A. & S.) Fr., N., R., W., S., F., galopus (Pers.) Fr., N., W., S., P., and var. alba Fl. Dan., R., and var. nigra Fl. Dan., W., epipterygia (Scop.) Fr., S., vulgaris (Pers.) Fr., W., stylobates (Pers.) Fr., N., W., F.

Omphalia fibula (Bull.) Fr., N., W., P., umbellifera (Linn.) Fr., R. Pleurotus corticatus Fr., W., acerosus Fr., C., applicatus (Batsch) Quél., S.

Hygrophorus conicus (Scop.) Fr., S., nigrescens Quél., P., chlorophanus Fr., S. Lactarius torminosus (Schaeff.) Fr., W., S., turpis (Weinm.) Fr., W., S., G., F., P., Lactarius torminosus (Schaeff.) Fr., W., S., turpis (Weinm.) Fr., W., S., G., F., P., blennius Fr., W., S., chrysorheus Fr., W., vellereus Fr., P., deliciosus (Linn.) Fr., W., pallidus (Pers.) Fr., R., W., quietus Fr., W., S., F., P., G., theiogalus (Fr.) Plowr., W., P., vietus Fr., W., S., rufus (Scop.) Fr., R., W., F., P., glyciosmus Fr., R., W., S., serifluus (DC.) Fr., N., S., P., mitissimus Fr., S., subdulcis (Pers.) Fr., W., S., tabidus Fr., W.

Russula chloroides (Krombh.) Bres., N., W., nigricans (Bull.) Fr., N., R., G., adusta (Pers.) Fr., R., W., S., P., G., azurea Bres., R., cyanoxantha (Schaeff.) Fr., N., R., W., S., G., furcata (Pers.) Fr., W., heterophylla Fr., W., S., P., G., pectinata (Bull.) Fr., N., ochroleuca (Pers.) Fr., N., R., W., S., P., G.

G., pectinata (Bull.) Fr., N., ochroleuca (Pers.) Fr., N., R., W., S., P., G., claroflava Grove, W., fellea Fr., N., R., W., S., P., drimeia Cooke, W., P., and var. Queletii (Fr.) Bat., N., fragilis (Pers.) Fr., W., S., P., emetica (Schaeff.) Fr., R., W., S., atropurpurea (Krombh.) Maire, N., S., P., G., xerampelina (Schaeff.) Fr., S., grisea (Pers.) Bres., R., puellaris Fr., W., vesca Fr., W., S., caerulea Cooke, W., sphagnophila Kauffm., S., W., paludosa Britz., W.

Cantharellus cibarius Fr., R., W., P., S., infundibuliformis (Scop.) Fr., W.
Marasmius peronatus (Bolt.) Fr., N., W., S., G., oreades (Bolt.) Fr., R., conigenus
(Pers.) Karst., R., W., G., erythropus (Pers.) Fr., W., S., P., hariolorum (DC.) Quél., N., S., P., dryophilus (Bull.) Karst., N., F.

Androsaceus androsaceus (Linn.) Pat., W., S., epiphyllus (Fr.) Pat., S.

Panus torulosus (Pers.) Fr., W., S. Volvaria bombycina (Schaeff.) Fr., S.

Pluteus cervinus (Schaeff.) Fr., N., P., S., salicinus (Pers.) Fr., W., pellitus (Pers.)

Entoloma Bloxamii Berk., B., sericeum (Bull.) Fr., S., B. Nolanea proletaria Fr., S., cetrata (Fr.) Schroet., R., S. Leptonia formosa Fr., B., sarcita (Fr.) Quél., S. Clitopilus prunulus (Scop.) Fr., W., S., P.

Claudopus variabilis (Pers.) W. G. Sm., S., G.

Paxillus involutus (Batsch) Fr., N., R., F., W., S., G., atrotomentosus (Batsch) Fr., W., giganteus (Sow.) Fr., F.

Pholiota erebia Fr., N., W., togularis (Bull.) Fr., N., R., praecox (Pers.) Fr., S., radicosa (Bull.) Fr., G., squarrosa (Mull.) Fr., P., spectabilis Fr., N., F., P., flammans Fr., N., marginata (Batsch) Fr., W.

Hebeloma sinuosum Fr., S., fastibile Fr., R., mesophaeum Fr., R., W., S., P., F., crustuliniforme (Bull.) Fr., S., P. Inocybe pyriodora (Pers.) Fr., Ś., tomentosa (Jungh.) Quél., W., corydalina Quél., W., geophylla (Sow.) Fr., R., W., P., G., and var. lilacina Fr., W., descissa Fr., S., obscura (Pers.) Fr., S., lacera Fr., W., S., cincinnata Fr., W., fastigiata

Astrosporina umbrina (Bres.) Rea, W., petiginosa (Fr.) Rea, W. Galera tenera (Schaeff.) Fr., W., S., spartea Fr., S., rubiginosa (Pers.) Fr., B., hypnorum (Schrank) Fr., N., S., F., P.

Naucoria semiorbicularis (Bull.) Fr., S., Myosotis Fr., W., sobria Fr., N., escharoides Fr., W., badia Kühner (=N. umbrina R. Maire), W.
Tubaria furfuracea (Pers.) W. G. Sm., W., S., crobulus Fr., S.
Flammula gummosa (Lasch) Fr., W., carbonaria Fr., P., fusus (Batsch) Fr., W., sapinea Fr., N., R., W., P., tricholoma (A. & S.) Fr., F., P., scamba Fr., P.

Cortinarius (Phlegmacium) varius (Schaeff.) Fr., W., largus Fr., S., multiformis Fr., S., caerulescens Fr., P., croceo-caeruleus (Pers.) Fr., S. (Myxacium) elatior Fr., R., W., S., (Inoloma) pholideus Fr., W., (Dermocybe) tabularis (Bull.) Fr., R., S., caninus Fr., R., S., anomalus Fr., S., phoeniceus (Bull.) Maire, S., semisanguineus (Brig.) Maire, W., cinnamomeus (Linn.) Fr., W., (Telamonia) bivelus Fr., S., hinnuleus (Sow.) Fr., S., P., hemitrichus Fr., W., rigidus (Scop.) Fr., S., P., (Hydrocybe) leucopus (Bull.) Fr., P., obtusus Fr., W., decipiens (Pers.) Fr., W.

Crepidotus mollis (Schaeff.) Fr., W.

Psalliota flavescens Gill., S., campestris (Linn.) Fr., N., sylvicola (Vitt.) Fr., N., S., haemorrhoidaria Kalchb., F., arvensis (Schaeff.) Fr., var. purpurascens Cooke, S., comtula Fr., W.

Stropharia aeruginosa (Curt.) Fr., N., W., P., G., squamosa (Pers.) Fr., S.,

coronilla (Bull.) Fr., C., semiglobata (Batsch) Fr., P.

Hypholoma sublateritium (Schaeff.) Fr., R., capnoides Fr., W., fasciculare (Huds.) Fr., N., R., W., F., P., G., epixanthum Fr., S., dispersum Fr., P., velutinum (Pers.) Fr., W., appendiculatum (Bull.) Fr., W., hydrophilum (Bull.) Fr., \mathcal{N} ., R., W., S.

Psilocybe sarcocephala Fr., R., ericaea (Pers.) Fr., W., subericaea Fr., W., uda (Pers.) Fr., B., P., semilanceata Fr., N., W., P., foenisecii (Pers.) Fr., W., C. Psathyra gossypina (Bull.) Fr., W., fibrillosa (Pers.) Fr., W., S.

Psathyrella gracilis Fr., W., crenata (Lasch) Fr., S., atomata Fr., C., F.

Panaeolus campanulatus (Linn.) Fr., N., W., F. Coprinus comatus (Fl. Dan.) Fr., F., atramentarius (Bull.) Fr., N., micaceus (Bull.) Fr., N., S., lagopus Fr., N., W., P., plicatilis (Curt.) Fr., C., ephemerus (Bull.)

Fr., S.
Gomphidius viscidus (Linn.) Fr., N., P.
Boletus luteus (Linn.) Fr., R., W., elegans (Schum.) Fr., W., P., viscidus (Linn.) Fr., W., granulatus (Linn.) Fr., N., badius Fr., W., S., bovinus (Linn.) Fr., W., piperatus (Bull.) Fr., W., variegatus (Sw.) Fr., R., W., chrysenteron (Bull.) Fr., N., S., F., P., G., subtomentosus (Linn.) Fr., S., N., F., versicolor Rostk., W., pruinatus Fr., R., edulis (Bull.) Fr., W., P., pinicola (Vitt.) Rea, P., reticulatus (Schaeff.) Boud., W., N., calopus Fr., W., erythropus (Pers.) Quél., N., R., P., W., S., duriusculus Schulz., W., versipellis Fr., R., S., G., scaber (Bull.) Fr., W., S., G.
Polyporus perennis (Linn.) Fr., W., P., G., brumalis (Pers.) Fr., S., squamosus (Huds.) Fr., N., Schweinitzii Fr., R., W., giganteus (Pers.) Fr., S., F., betulinus (Bull.) Fr., R., W., P., benzoinus (Wahlenb.) Fr., N., adustus (Willd.) Fr., R.,

(Bull.) Fr., R., W., P., benzoinus (Wahlenb.) Fr., N., adustus (Willd.) Fr., R., W., S., lacteus Fr., R., fragilis Fr., R., W., caesius (Schrad.) Fr., R., stipticus

(Pers.) Fr., N., S.

Fomes annosus Fr., N., R., W., S., F., P. (on birch).

Ganoderma applanatum (Pers.) Pat., N., S.

Poria sanguinolenta (A. & S.) Fr., R., hymenocystis B. & Br., N., xantha (Fr.) Lind, P.

Polystictus versicolor (Linn.) Fr., N., W., abietinus (Dicks.) Fr., N.

Irpex obliquus (Schrad.) Fr., F., S., G.

Lenzites betulina (Linn.) Fr., W.

Trametes gibbosa (Pers.) Fr., W., rubescens (A. & S.) Fr., on Salix, G.

Merulius tremellosus (Schrad.) Fr., W., P.

Phlebia merismoides Fr., R.

Fistulina hepatica (Huds.) Fr., N.

Hydnum repandum (Linn.) Fr., W., G., auriscalpium (Linn.) Fr., R., W., P. Mycoleptodon ochraceum (Pers.) Pat., S.

Acia uda (Fr.) Bourd. & Galz., S.

Grandinia farinacea (Pers.) Bourd. & Galz. S., granulosa Fr., S.

Odontia arguta (Fr.) Quél., R., papillosa (Fr.) Bres., R. Thelephora terrestris Ehrh. ex Fr., N., R., W., F., P., G.

Hypochnus fuscus (Pers.) Fr., S., F., fumosus Fr., R., W., F.

Sparassis crispa (Wulf.) Fr., W., S., P.

Stereum spadiceum Fr., P., hirsutum (Willd.) Fr., N., W., P., rugosum (Pers.) Fr., R., W., P., purpureum Pers. ex Fr., S., P.

Hymenochaete Mougeotii (Fr.) Cooke, N., R.
Corticium fuciforme (Berk.) Wakef., N., W., arachnoideum Berk., R., Sambuci (Pers.) Fr., N., subcoronatum v. H. & Litsch., R., confine Bourd. & Galz., R., S., praetermissum (Karst.) Bres., S.

Peniophora byssoidea (Pers.) v. H. & Litsch., W., velutina (DC.) Cooke, S., setigera (Fr.) Bres., S., hydnoides Cooke & Mass., R., S., gigantea (Fr.) Mass., N., W., incarnata (Pers.) Cooke, S., W., quercina (Pers.) Cooke, N., P.

Cyphella capula (Holmsk.) Fr., S. Solenia anomala (Pers.) Fr., N.

Clavaria cristata (Holmsk.) Fr., R., cinerea (Bull.) Fr., S., rugosa (Bull.) Fr., R., W., inaequalis (Mull.) Fr., N., fistulosa (Holmsk.) Fr., S., R., W. Pistillaria quisquiliaris Fr., R., S., puberula Berk., W., S.

Auricularia auricula-Judae (Linn.) Schroet., R.

Exidia glandulosa (Bull.) Fr., S.

Tremellodon gelatinosum (Scop.) Pers., R., S. Sebacina incrustans (Pers.) Tul., P., S. Dacryomyces deliquescens (Bull.) Duby, N., W., S.

Calocera viscosa (Pers.) Fr., R., S., F., P., cornea (Batsch) Fr., R., stricta Fr., R.

GASTEROMYCETES

Cynophallus caninus (Huds.) Fr., W., P., S.

Phallus impudicus (Linn.) Pers., N., R., P., S.
Lycoperdon saccatum (Vahl) Fr., W., S., umbrinum Pers., W., S., perlatum Pers.,
W., S., N., pyriforme (Schaeff.) Pers., S., P.
Bovistella paludosa (Lév.) Lloyd, B.

Geaster triplex Jungh., S. Crucibulum vulgare Tul., R., S., F. Cyathus striatus (Huds.) Pers., S.

Scleroderma aurantium Pers., N., W., P. Sphaerobolus stellatus (Tode) Pers., R., S., P.

UREDINALES

Uromyces Rumicis (Schum.) Wint., N., G., Acetosae Schroet., N., Valerianae (Schum.) Fuck., B., G.

Puccinia calthaecola Schroet. on Caltha palustris, B., G., Thalictri Chev., G., Violae (Schum.) DC., W., P., aegra Grove, F., G., Lychnidearum Link, N., Malvacearum Mont., C., Cicutae Lasch, G., Hydrocotyles (Link) Cooke, W., Saniculae Grev., S., Centaureae Mart., N., obtegens (Link) Tul., F., Hypochaeridis Oud., B., Antirrhini Diet. & Holw., F., C., Glechomatis DC., N., C., P., Menthae Pers., R., G., annularis (Str.) Schlecht., N., R., W., B., Polygoni A. & S., S., Buxi DC., N., Caricis (Schum.) Rebent., G., Lolii Niels. C., Magnusiana Körn., G., Phragmitis (Schum.) Körn., G., Poarum Niels., aecidium on Tussilago, S., mirabilissima Peck, F., P.

Phragmidium violaceum (Schultz.) Wint., N., W., Rubi (Pers.) Wint., G.

Kuehneola albida (Kuehn.) Magn., N., W., F., Tormentillae (Fuck.) Arth., W., B.

Coleosporium Euphrasiae (Schum.) Wint., B., Senecionis (Pers.) Fr., W., C., Tussilaginis (Pers.) Kleb., S.

Cronartium asclepiadeum Fr. on Tropaeolum majus, F.

Pucciniastrum Agrimoniae (DC.) Tranzsch., S., G., Epilobii (Pers.) Otth, W., R., S., P., F.

Melampsoridium betulinum (Pers.) Kleb., W., R., P., S. Milesina Blechni Syd., W.

USTILAGINALES

Ustilago Lychnidis-dioicae (DC.) Liro, N., R., longissima (Schlecht.) Meyen, G., utriculosa (Nees) Tul. on Polygonum hydropiper, G. Sphacelotheca Hydropiperidis (Schum.) de Bary, C. Tilletia striaeformis (West.) Wint., P.

PLECTASCALES

Ctenomyces serratus Eidam, R. Elaphomyces granulatus Fr., W.

PYRENOMYCETES

Sphaerotheca Humuli (DC.) Burr. on Arctium, P., on Epilobium, G., pannosa (Wallr.) Lév., N.

Erysiphe Polygoni DC. on Delphinium, \mathcal{N} ., on Swede, \mathcal{N} .

Microsphaera Grossulariae (Wallr.) Lév., P.

Uncinula Aceris (DC.) Sacc., R., P.

Phyllactinia corylea (Pers.) Karst. on Corylus, W., P. Gibberella pulicaris (Fr.) Sacc. on Broom, Mousehold Heath.

Nectria cinnabarina (Tode) Fr., conidial only, on Ilex, W., on Ulex, S., Desmazierii de Not. on Buxus, N., Aquifolii (Fr.) Berk. on Ilex, S., punicea (Kunze & Schm.) Fr. on Ilex, S.

Hyponectria Buxi (Desm.) Sacc. on Buxus, \mathcal{N} .

Hypocrea pulvinata Fuck. on Polyporus betulinus, P. Claviceps microcephala (Wallr.) Tul., S.

Cordyceps ophioglossoides (Ehrh.) Link, W., capitata (Holmsk.) Link, R., W. Lasiosphaeria hirsuta (Fr.) Ces. & de Not., G. Leptospora ovina (Pers.) Fuck., S.

Diaporthe eres Nits. on Ulex, S., fibrosa (Pers.) Fuck. on Rhamnus cathartica, G., leiphaemia (Fr.) Sacc. on Quercus, S.

Peroneutypa heteracantha (Sacc.) Berl. on Acer, N.

Cryptosphaeria eunomia (Fr.) Fuck. on Fraxinus, G.
Melanconis stilbostoma (Fr.) Tul. on Betula, W.
Pseudovalsa lanciformis (Fr.) Ces. & de Not. on Betula, R.
Diatrype disciformis (Hoffm.) Fr. on Acer, N., on Carpinus, S., Stigma (Hoffm.) Fr. on Carpinus, S.

Hypoxylon coccineum Bull. on Fagus, W., argillaceum (Pers.) Fr. on Fraxinus, G., multiforme Fr. on Pyrus Aucuparia, S., semiimmersum Nits. on Quercus, W.

Daldinia concentrica (Bolt.) Ces. & de Not., F.

Xylaria Hypoxylon (Linn.) Grev., W., S., P., polymorpha (Pers.) Grev., N., S., P., Tulasnei Nits. on rabbit dung, W.

Phyllachora graminis (Pers.) Fuck. on Dactylis, N.

Rhopographus filicinus (Fr.) Nits., N., P.

HYSTERIALES

Gloniopsis curvata Sacc. on Fraxinus, G. Hysterium angustatum A. & S. on Fraxinus, G.

DISCOMYCETES

Helvella crispa (Scop.) Fr., R., W., S., F., lacunosa Afz., S. Macropodia macropus (Pers.) Fuck., W. Rhizina inflata (Schaeff.) Karst., P. Galactinia badia (Pers.) Boud., W., P. Otidea onotica (Pers.) Fuck., N. Geopyxis carbonaria (A. & S.) Sacc., P. Peziza aurantia Pers., N., R.

Pulvinula constellatio (Cooke) Boud., S. Saccobolus violascens Boud., S. Taphrina Tosquinetii (West.) Magn., W. Leotia lubrica (Scop.) Pers., W. Cudoniella acicularis (Bull.) Schroet., R., S. Coryne sarcoides (Jacq.) Tul., W., S. Bulgaria inquinans (Pers.) Fr., N., P. Corynella atrovirens (Pers.) Boud., S. Orbilia xanthostigma Fr., W. Phialea echinophila (Bull.) Quél., S., firma (Pers.) Gill., S. Chlorosplenium aeruginosum (Oeder) de Not., S., on beech. Helotium fructigenum (Bull.) Fuck., R., S. Trichoscypha calycina (Schum.) Boud., W. Mollisia cinerea (Batsch) Karst., S., P. Phacidium multivalve (DC.) Kunze & Schm., W. Stegia Ilicis Fr., W., S. Rhytisma acerinum (Pers.) Fr., N., F., P.

PHYCOMYCETES

Syzygites megalocarpus Ehrenb., W., S. Bremia Lactucae Regel, S. Plasmopara nivea (Unger) Schroet., on Aegopodium, P. Peronospora alta Fuck., on Plantago, F. Entomophthora echinospora Thaxter, on fly, P.

DEUTEROMYCETES

Dendrophoma pruinosa (Fr.) Sacc. on Fraxinus, G. Actinonema Rosae (Lib.) Fr., N. Eleutheromyces subulatus (Tode) Fuck. on Sparassis, W. Septoria Violae West. on Viola Riviniana, W. Cylindrium flavovirens Bon., R. Trichoderma viride Fr., N. Rhinotrichum Thwaitesii B. & Br., W., P. Sepedonium chrysospermum (Bull.) Fr., N., R. Ovularia obliqua (Cooke) Oud., S., P., haplospora (Speg.) Magn. on Alchemilla arvensis, N. Botrytis cinerea Pers., R. Cephalosporium coccorum Petch, on Mealy Bugs, S. Acremonium album Preuss on Stemonitis, S. Gliocladium penicillioides Corda on Gibberella, Mousehold Heath. Chalara fungorum Sacc. on Eleutheromyces, W.

Ramularia Urticae Ces., W., acris Lindr. on Ranunculus acris, P., Tulasnei Sacc., S., calcea (Desm.) Ces., N., R., F., P., sambucina Sacc., N., P., Primulae

Napicladium arundinaceum (Corda) Sacc., on Phragmites, G. Stilbella erythrocephala (Ditm.) Lindau, N., F., W.

Tilachlidium tomentosum (Schrad.) Lindau, on Cribraria, R., W., S., on Comatricha, R.

Isaria (Tilachlidium) brachiata (Batsch) Schum., on decaying Agaric, R., (Spicaria)

farinosa (Holmsk.) Fr., on pupae, R., W., P. Gibellula aranearum (Schw.) Syd. on spider, P.

Sporocybe Azaleae Peck, N., W. Tuberculina persicina (Ditm.) Sacc., on Puccinia Poarum, S., on P. Antirrhini, Norwich.

Volutella Buxi (Corda) Berk., P. Anthina flammea Fr., S.

Ectostroma Iridis (Ehrenb.) Fr. on Iris pseudacorus, G.

MYCETOZOA FOUND DURING THE NORWICH FORAY

By H. J. HOWARD

The summer had been characterised by conditions unfavourable to the development of Mycetozoa, but a week's rain previous to the foray and rain at the end of almost every day caused several species to appear and a total of thirty-six was recorded.

October 2nd. A visit was made to Northrepps Hall Woods, which consist chiefly of beech with very few old logs: nothing of outstanding importance was gathered. The afternoon, spent in West Runton woods where a number of pine logs were present, produced *Tubifera ferru*-

ginosa and Cribraria vulgaris.

October 3rd was a dull day and the extensive Westwick woods were visited: these consist of heathland, with pine wood and Sphagnum under birches. It is probable that it was on this day that the new British record of Physarum javanicum was made by Miss Cayley, who collected this species on bark with Parmelia sp. Owing to its superficial resemblance to Physarum nutans it was taken for that species, but on being sent to Miss Lister the discoid sporangia with the deposits of lime uniformly distributed in the walls and not clustered to form spots as in all varieties of P. nutans led to its identification. Later in the afternoon the North Walsham woods were explored in the same neighbourhood.

October 4th. Stratton Strawless woods, with mixed growth of beech, birch, hornbeam, oak, chestnut, bramble and bracken, with numerous clearings, yielded a variety of Mycetozoa. The journey was continued to Buxton Heath, but a rain storm prevented extensive

searching there.

October 5th. Sprowston and Plumstead Road Woods and Framingham Chase proved somewhat unfruitful. The afternoon, however, provided an interesting diversion, for a chance meeting with Mr J. Morse of Eaton resulted in a visit to his cucumber-houses where *Physarum gyrosum* was seen in abundance, both in the plasmodium and fruiting stages. Sticks, soil and stems of cucumbers were covered with rosettes of this species and sometimes the considerable patches of plasmodium had crept up the pots standing upon the soil. This species has occurred in these glasshouses in great abundance on several occasions and, it is interesting to note, always in houses manured with sewage sludge and never in others adjacent treated with farmyard manure. A piece of soil covered with creamy white veins of plasmodium was taken home which next day was found to have changed

to "clear Amazonite blue" as described by M. K. Minakata from Tanabe, Japan. Unfortunately instead of being allowed to mature naturally, the mass was placed in a very moist incubator in order to imitate the conditions of the cucumber house, and the temperature being too high, the plasmodium was injured and turned to a bloodred colour in drying. I am inclined to think that the plasmodium normally changes to blue before forming sporangia, but hope to make confirmatory observations should the species appear again. Fuligo septica var. candida was also found maturing upon cucumber stems.

 \mathcal{N} . = Northrepps Hall Woods; R. = West Runton Woods; W. = Westwick and North Walsham Woods; S. = Stratton Strawless Woods; P. = Sprowston and Plumstead Road Woods; F. = Framingham Chase; E. = cucumber houses at Eaton, Norwich.

Ceratiomyxa fruticulosa (Muell.) Macbr., N., pine wood.

Badhamia utricularis Berk., S., dead pine wood. Physarum nutans Pers., N., W., dead wood, var. leucophaeum Lister, S., F., dead wood.

P. javanicum Racib., W.?, bark with lichen.

P. gyrosum Rost., E., soil, stems, pots, in cucumber houses.

P. bitectum Lister, W., old larch.

Fuligo septica (L.) Gmel., N., R., W., F., dead wood, var. candida Lister, N., pine wood, E., cucumber stems.

Craterium minutum (Leers) Fr., W., S., dead leaves, etc. C. leucocephalum Ditm., W., holly leaves.

Leocarpus fragilis (Dicks.) Rost., W., S., beech leaves, twigs.

Diderma hemisphericum (Bull.) Hornem., W., twigs and alder leaves.

Diachaea leucopoda Rost., S., twigs and leaves. Didymium difforme (Pers.) Duby, W., dead leaves. D. Clavus (A. & S.) Rost., N., S., dead leaves. D. melanospermum (Pers.) Macbr., W.

D. nigripes Fr. var. xanthopus Lister, W., S., alder twigs, etc. Mucilago spongiosa (Leysser) Morg., W., S., grass. Stemonitis fusca Roth, W., S., dead wood. S. splendens Rost. var. flaccida Lister, W., alder.

S. herbatica Peck, R., twigs and leaves.

S. flavogenita Jahn, S., P.

Comatricha nigra (Pers.) Schroet., W., S., P., birch, etc. C. typhoides (Bull.) Rost., S., dead wood.

Enerthenema papillatum (Pers.) Rost., W., S., P., dead wood.

Cribraria argillacea Pers., S., dead wood.

C. rufa (Roth) Rost., R., pine wood.

C. vulgaris Schrad., R., W., S., F., pine wood, var. aurantiaca Pers., R., S., pine wood.

Licea flexuosa Pers., N., S., green slimy pine log, etc.
Tubifera ferruginosa Gmel., N., R., W., S., F., dead pine wood.
Lycogala epidendrum Fr., E., twigs in cucumber house.
Trichia persimilis Karst., N., dead wood.

T. varia Pers., W., S., dead birch, etc.

Arcyria cinerea (Bull.) Pers., W., F., dead wood.

A. denudata Wettst., W., S., birch, etc. A. incarnata Pers., N., R., W., S., dead wood. A. nutans (Bull.) Grev., R., dead wood.

NORFOLK LICHENS

By H. H. KNIGHT

The Norfolk woods visited during the Foray were not very rich in lichens, and in the woods near Norwich visited on Friday, particularly those of Sprowston and Plumstead, the trees were very bare. Graphis elegans and scripta were seen in Stratton Strawless woods, but nowhere else, and no species of Opegrapha was seen on trees. Lecidea lucida occurred on a brick wall in the woods near Overstrand. This is a lichen which appears to be spreading from its natural habitat on siliceous rocks to brick walls, and is sometimes found even on town walls in the Midlands.

On October 6th I visited the Marams near Blakeney with Dr Watson. On the stones here a number of saxicolous lichens were found. Physcia ciliaris, usually a tree lichen, was growing on the ground, and *Placodium luteoalbum*, which prefers elms, was growing on the stems of Suaeda fruticosa.

In the following list I have as usual followed the order and naming of the Monograph of the British Lichens, by Miss A. Lorrain Smith. Without the help of Dr Watson the list would have been much shorter.

O. = Overstrand and Runton Woods; W. = Westwick and North Walsham Woods; S. = Stratton Strawless Woods; B. = Walls and trees near Woodrow Inn and Buxton Heath; P. = Plumstead and Sprowston Woods; F. = Framingham Woods; M. = The Marams near Blakeney and walls near Cley; C. = Common lichens.

Chaenotheca melanophaea Zwackh.,

Calicium hyperellum Ach., S., M. Cyphelium inquinans Trev., O. Peltigera canina Willd., O., W., M.

P. polydactyla Hoffm., W., F. Parmelia physodes Ach., C. var. platyphylla Ach., F.

P. perlata Ach., O., S. P. caperata Ach., W., S., F. P. subaurifera Nyl., S.

P. sulcata Tayl., C.

P. dubia Tayl., S.

P. revoluta Floerke, W., S. P. acetabulum Dub., F.

P. fuliginosa Nyl., F., M. var. laetevirens Nyl., C.

Cetraria aculeata Fr., M. Evernia prunastri Ach., C. Ramalina calicaris Fr., S.

R. fastigiata Ach., S.

Ramalina farinacea Ach., C.

R. pollinaria Ach. f. humilis Cromb.,

Xanthoria parietina Th. Fr., C.

var. aureola Th. Fr., S., M. X. polycarpa Oliv., W., M. X. lychnea Th. Fr., B., F.

Placodium flavescens A. L. Sm., M.

P. murorum DC., B., M. var. pusillum Flag., M.

P. lobulatum A. L. Sm., M. P. citrinum Hepp, B., M.

P. phloginum A. L. Sm., S.

P. aurantiacum var. flavovirescens Hepp, M.

P. luteoalbum Hepp, M.

P. atroflavum A. L. Sm., M. Candelariella vitellina Müll.-Arg., C.

Physcia ciliaris DC., M. P. pulverulenta Nyl., M.

P. grisea A. Zahlbr., S.

Physcia hispida Tuckerm., S., M. P. caesia Nyl., B. P. orbicularis var. virella Dalla Torre & Sarnth., B., M. Rinodina demissa Arn., *M.* R. umbrinofusca Oliv., M. Lecanora muralis Schaer., B. L. subfusca Ach., O., P. var. chlarona Ach., S., F. var. allophana Ach., S. L. rugosa Nyl., S. L. intumescens Koerb., P. L. campestris B. de Lesd., B., M. L. atra Ach., *M*. L. Hageni Ach., S., F., M. L. umbrina Massal., M. L. galactina Ach., B., M. L. dispersa Nyl., M. L. varia Ach., C. L. conizaea Nyl., F. var. conizaeoides A. L. Sm., S., P., F. L. symmictera Nyl., B., F., M. L. expallens Ach., S. Lecania prosechoides A. L. Sm., M. L. erysibe Mudd, M. Pertusaria globulifera Nyl., S. P. faginea Leight., C. P. pertusa Dalla Torre & Sarnth., C. Phlyctis agelaea Koerb., S., F. Baeomyces rufus DC., W., F. Cladonia sylvatica Hoffm., O., B. C. foliosa Willd., M. C. pyxidata Hoffm., C. var. chlorophaea Floerke, S., F. C. fimbriata Fr., C. var. simplex Wain., F. var. radiata Cromb., S., F. var. subcornuta Nyl., F. C. ochrochlora f. ceratodes Floerke, B. C. pityrea Fr., B. C. crispata Flot., B.

Cladonia furcata Schrad., C. var. spinosa Leight., M. var. recurva Hoffm., B. C. rangiformis Hoffm., M. var. foliosa Wain., M. C. squamosa Hoffm., S. C. digitata Hoffm., W., F. C. coccifera Willd., B. C. flabelliformis Wain., F. C. macilenta Hoffm., B., F. C. Floerkeana Fr. var. carcata Wain., B. Gyalecta diluta Wain., P., F. Lecidea lucida Ach., O. L. coarctata Nyl., F. L. quernea Ach., S. L. granulosa Schaer., C. L. flexuosa Nyl., W. L. uliginosa Ach., S. L. fuliginea Ach., S. L. dubia Hook., M. L. expansa Nyl., M. Biatorina Griffithii Massal., S., P., F. B. erysiboides Th. Fr., F. B. Lightfootii var. commutata Mudd, S. B. lenticularis Koerb., M. B. chalybeia Mudd, M. Bacidia phacodes Mudd, S. Buellia canescens De Not., B., M. B. myriocarpa Mudd, B., M. B. stellulata Mudd, M. B. confervoides Kremp., M. Rhizocarpon confervoides DC., M. Graphis elegans Ach., S. G. scripta Ach., S. Verrucaria maura Wahlenb., M. V. microspora Nyl., M. V. viridula Ach., B., M. V. nigrescens Pers., M. V. mutabilis Borr., F. V. muralis Ach., F. Acrocordia epipolaea A. L. Sm., B.

PRESIDENTIAL ADDRESS

By B. BARNES, D.Sc., Ph.D., F.L.S.

INDUCED VARIATION

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m P}_{
m ROBABLY}$ ever since man has concerned himself with domesticated plants and animals, he has tried to improve his stocks. Much success has been attained by judicious selection and by systematic breeding from chance variations of a desirable kind which turned up in the animals and plants, and efforts have been made by artificial means to provoke variations. There is a traditional method of obtaining variation by the simultaneous use of high feeding, relatively high temperatures and crowding; this method seems to have been effective in dealing with cultivated plants, and the Chinese are said to have brought about much alteration in gold fish by keeping them in confined spaces in dirty conditions. Darwin, in his Variation under Domestication,* refers to the general opinion among plant breeders that a stock of plants must be grown on poor land if it is to be kept true. Such ideas, widely held by practical men whose living depends on success with their plants and animals, must be based on a good deal of experience, and cannot be disregarded, but the traditional methods appear to be slow and capricious in their operation, and do not seem capable of much control.

In the second half of the nineteenth century, some scattered work was done on the experimental induction of variation, but at that time there was little sympathy with any work which tended to upset the idea of the fixity of species, a somewhat strange position, since variability, and very great variability, was accepted without difficulty in domesticated creatures. A growing realisation that experiment was necessary in order to get a better understanding of the mechanism of evolution, and the development of Mendelism, combined to awaken interest in variation, and stimulated efforts to discover if the rate of change in organisms could be influenced by direct treatment with physical or chemical agents, since the successful outcome of such experiments might well throw light on the nature of evolutionary changes, and also free the breeder from the necessity of waiting until some chance variant fell into his hands. Up to the present, we have not made much progress in elucidating the mysteries of evolution, and it is not clear that the breeder has been much helped, but a number of workers, using many kinds of plants and animals, have shown, during the present century, that changes can be induced by the use of chemicals, by the application of high tem-

^{*} Darwin, C., Variation under Domestication, II (1905), 300-3. London: Murray.

peratures, by dosage with X-rays and with emanations from radium, by exposure to ultra-violet light, and, probably, by taking advantage of some slow changes going on within the organisms. There is no evidence at present that any of the agents found to be effective have any specific effect; similar variations have been induced in the same stock of the same organism after treatment with high temperatures or after exposure to X-rays. The experimental results have been and continue to be erratic, since the work is difficult, and since, so far, no one has invented a technique capable of exact application. Even with such simple things as bacteria and fungal spores, it is impossible to be sure that the material is homogeneous and that the treatment is evenly applied; with seeds of the higher plants, with growing plants, and with the eggs, and other developmental stages of animals, standardisation of the experimental material is impossible. Although as yet we have no means of repeating an experiment with a reasonable expectation of repeating the results of previous similar experiments, the failure of one worker to repeat exactly the work of another

does not mean that the results of either are valueless.

The simple organisation of the fungi, and the ease with which they are grown in pure culture, makes these organisms very suitable subjects for experiments on induced variation, and fungi have afforded as satisfactory evidence as have any group of organisms that variation can be induced. Hansen,* well known for his work on yeasts, produced the first good evidence of the induction of variation in a fungus. He developed a sound technique for the isolation of single cells, and for the maintenance of pure cultures started from single cells. After he had shown that yeasts continued to bud at a temperature a few degrees higher than the maximum temperature at which they could form spores, Hansen succeeded in producing a non-sporing race from a richly sporing stock of a wine yeast (Johannisberg II of Wortmann, probably a form of Saccharomyces ellipsoideus) by prolonged culture at a temperature between the maximum for spore formation and the maximum for budding. The original material lost the power of sporulation gradually, but, after working for about three months with cultures which were renewed every few days, he obtained a race which did not form spores, no matter how it was fed or treated; this race was kept in culture for sixteen years, and during that time did not regain the power of sporulation. Since the original yeast was distinguished by its rich sporulation, and since Hansen could never show that it produced non-sporing cells in ordinary culture, he fairly concluded that the asporogenous race had come into existence because his experiments had caused a change, and that it had not been grown from a non-sporing cell present in the original material, and accidentally selected as a starting point for the cultures.

^{*} Hansen, E. C., C.R. Lab. Carlsberg, 11 (1883), ii; Ann. Bot. 1x (1895), 549.

Aspergillus niger has long been a favourite subject for experiments with fungi; a Continental writer once called it the mycologist's guinea-pig. A. niger has yielded induced variants after exposure to high temperatures, and after treatment with chemicals. In 1912, Schiemann* gave a very thorough account of changes produced in a stock of the fungus, a stock well known to be constant in ordinary culture. She distinguished carefully between modifications due to the use of special conditions, and only appearing so long as the special conditions were operative, and true permanent changes. For example, media containing copper sulphate in a concentration of 1/1000, and known to be of a kind very favourable to the growth of the fungus when copper sulphate was not present, bore weak whitish colonies with poor crops of conidia; these weak colonies were however nothing more than modifications, for, conidia taken from them and planted on ordinary media yielded the normal A. niger at once. On the other hand, the addition of potassium bichromate to the medium sometimes caused the appearance of a true variant. It was found that although some growth was possible on media containing potassium bichromate in a concentration greater than 1/2000, such concentrations prevented the forming of conidia. On media whose content of the salt was just below that necessary to prevent sporulation, germination was greatly delayed (taking sixteen days instead of about a day), and fruiting was tardy (needing twenty-eight days instead of three to four). One of the first cultures made on a medium of this kind yielded some rusty brown conidial heads among the normal blackish heads, and the light-coloured heads yielded a variant which was still alive in culture in 1929, and may be so still, developing its light-coloured heads on ordinary media, free from bichromate. The early isolations of the variant, on ordinary media, showed colour fluctuations about the average chocolate-brown, but these fluctuations were inconstant and could not be transmitted to further cultures. Such initial instability is a common feature when variants are settling down after their first appearance.

A cinnamon-brown variant appeared after A. niger had been passed through eleven successive cultures on a medium containing potassium bichromate in a concentration of 1/20,000; the early isolations were unstable, and went through several transfers before settling into a

permanent variant.

Schiemann also tried the effects of high temperatures. When A. niger was grown on an ordinary medium at 40–45° C., the growth was specially dense, the conidiophores were dwarfed, and other irregularities were noted; transfers from these cultures to an ordinary medium kept at 36° C., a temperature specially favourable to the growth of the fungus, always gave the normal form. Some cultures

^{*} Schiemann, E., Zeitschr. f. induk. Abstamm. u. Vererb. VIII (1912), 1.

were heated for a time to 48° C., and this treatment led to the appearance of true variants. From one of these cultures, a form was isolated showing characters the reverse of those shown by the high-temperature modifications of the species, for the variant grew loosely, produced conidiophores two to three times as long as normal conidiophores, and bore conidial heads slightly larger than normal heads. It follows that modifications due to some special treatment are not necessarily steps towards the formation of a permanent variant under

the influence of similar treatment.

Another heated culture produced a greyish head of conidia, and these conidia, planted out separately, yielded some colonies in which sectors appeared, the sectors fruiting later than the rest of the colony. Isolations from the sectors gave a form whose behaviour was inconstant; the first colonies grew weakly and irregularly, but as successive cultures were made, the fungus slowly gained strength. The colonies of this variant were very sensitive to slight differences in the composition and the moisture content of the medium, giving modifications readily, and, in this respect, the variant was more responsive than the stock strain of A. niger. The variant germinated and grew well at temperatures too low for the best growth of the parent strain, and was always late in sporing. Young colonies produced greyish brown conidial heads, but on older colonies pigmentation was usually normal. Transfers from young colonies gave the unstable variant; transfers from old colonies gave normal A. niger. These phenomena are of special interest; they suggest that the variant was not much altered, and that it had suffered a mild amount of change from the heating, so mild that recovery was possible after a mycelium had grown for a time. Abnormalities in youth followed by an apparent resumption of normal features at a later stage have been encountered in the induced variants of other organisms.

Schiemann's work shows clearly that a given species may be made to yield a series of forms ranging from modifications depending solely for their appearance on special conditions of culture, to pronounced

variants with a great degree of permanence.

Waterman* encountered variation in A. niger during an attempt to discover substances which were specially good sources of carbon for the fungus. One such substance, para-oxybenzoic acid, used in conjunction with the necessary mineral salts, was found at first to serve the fungus well, and to lead to the storage of much glycogen within the hyphae. Discrepancies began to appear in the chemical determinations however, and these were traced to the formation of variants under the influence of the para-oxybenzoic acid. A. niger was also found to vary after treatment with substances which checked its growth, so that both substances apparently favourable to growth and

* Waterman, H. J., Z. GärPhysiol. ni (1913), 1.

substances certainly unfavourable to growth led to the development of variants. An example such as this indicates the complexity of the position and suggests that the mechanism underlying induced variation can hardly be reduced to one universal explanation.

Haenicke* isolated light-coloured forms of A. niger from cultures which had been kept at 45° C.; one was accidentally killed after it had lasted through nine transfers, the other suddenly reverted to normal after retaining its aberrant characters through eighteen suc-

cessive cultures.

Between 1926 and 1932, many experiments were made† on the effect of high temperatures on Eurotium herbariorum, Botrytis cinerea and Thamnidium elegans. Well established strains of the three species were available, all known to be constant under ordinary conditions of culture. Spores taken from pure cultures, with precautions to avoid contamination, were heated in various ways and then planted on the usual media; at the same time unheated spores were planted on other dishes of the same media. The experiments yielded a considerable number of variants from the heated spores, while none was seen in the cultures from unheated spores. Since, during some years of culture, the three stocks had never thrown aberrant forms, it seems a fair conclusion that the development of variants in cultures started from heated spores must be due to the treatment given to the spores. Eurotium herbariorum responded best to treatment, no doubt because it offered the most possibilities for change, for the strain used readily formed conidia and perithecia in culture. Some of the variants were slight, and apparent only in young colonies; most of the slight variants reverted sooner or later to the normal form, persisting even in young colonies only through a few transfers. At the other end of the scale, some of the variants obtained in 1926 were still in existence, unchanged, in 1933, having kept their characters through more than 140 transfers made on ordinary media and started from unheated spores. The changes affected the rate of growth, the colour and morphology of the conidia and conidiophores, the colour, form and abundance of the perithecia, the form of the ascospores, and indeed every obvious morphological character of the fungus was changed in one variant or another; though the matter was not investigated systematically, variations in staling capacity, and variations in the extent to which the medium was stained, suggested that physiological characters had also been altered. Some variants were slightly unstable, with a tendency to sectoring, and, in one, sectoring was certainly stimulated by a change of medium. Isolations from these sectors gave another variant, evidently a weakened version of the first; it remained

^{*} Haenicke, A., Z. Bot. VIII (1916), 225. † Barnes, B., Ann. Bot. XLII (1928), 783; XLIV (1930), 825; Trans. Brit. mycol. Soc. XIX (1935), 291.

distinctfrom its parent for some eighteen months, but slowly reverted to the form of the original variant, but not to that of the stock Eurotium herbariorum. A brown variant was particularly distinguished by the heavy production of a sterile, white aerial mycelium on old colonies. Another, with conidia not greatly different in colour from those of the normal form, always showed some deformations of the conidial apparatus and never developed perithecia; this variant, and another which formed but few perithecia, stained the medium heavily. It was not possible to demonstrate that there was any general tendency in the variants for characters to be changed in groups, though variants which were able to develop good crops of perithecia seldom produced much aerial mycelium, and they also showed little tendency to stale, or to stain the medium.

Botrytis cinerea offered less possibility of change; alterations were remarked in the general habit, in the form, colour and abundance of the sclerotia, and in the morphology of the conidial apparatus. Some of the variants were extremely unstable when first isolated, and one notable variant was sterile, very weak, and so deficient in strength that it died out after a few transfers. Other variants remained recognisable for more than two years, gradually reverting to normal as time went on, and still others, in particular one that formed dense white colonies, appeared to be permanent. Staining of the medium and the production of much whitish aerial mycelium was most apparent in the variants which developed few or no sclerotia, and if the reasonable assumption be made that the sclerotia of Botrytis have some relation to sexual phenomena, we then have in these variants a parallel with some variants of Eurotium herbariorum.

Variants of *Thamnidium elegans* were obtained from heated sporangiospores, and they too showed a range from transient to apparently permanent forms. Partial sterility and increase of aerial mycelium characterised the more stable variants, but despite very marked alterations in morphology, all attempts to demonstrate any effect on the sexual reactions, by growing the variants in contrasted cultures, were fruitless; thus, in Zygomycetes, a difference in morphology between two strains of a heterothallic species does not necessarily

indicate a difference in sexual character.

An attempt was made during the investigation of the three species just discussed, to establish a relation between the severity of the initial heating and the frequency of variation, and between the severity of treatment and the degree of alteration. Conclusive evidence was not obtained, for many thousands of experiments would be needed to settle the point, but it appeared that exposure to heat just insufficient to kill the spores was most likely to produce the greatest number of striking alterations.

Experiments with heated ascospores of Eurotium herbariorum did not

succeed. Bean and Brooks* were unable to cause variation in *Pyronema confluens* by heating the ascospores, and Dickson† had the same experience with the ascospores of *Chaetomium cochliodes*, though he found that these spores produced many variants after treatment

with X-rays.

Irradiation with X-rays has provoked variation in many organisms, including fungi; as usual, the results of the experiments have been erratic. Nadson and Philippov‡ irradiated young mycelia of several species of Zygomycetes, with noteworthy effect; disturbances were noted in the amount of the crops of sporangia and zygospores, and in pigmentation. Isolations from treated mycelia of Mucor genevensis yielded sectoring colonies, and cultures from the sectors gave variants of greater or less permanency. An interesting variant of M. genevensis was first noted as a sector bearing very few zygospores and having globules of reddish yellow oil in the hyphae. Subsequently, a similar variant was obtained from irradiated material of Zygorhynchus Moelleri; it formed no zygospores, developed a heavy crop of sporangia, and showed the yellow hyphal inclusions. The connection between disturbances of fertility and the development of pigmentation suggests a parallel with fungi already considered, and further, the induction of reduced fertility and yellow pigmentation in Zygomycetes is of interest, since these characters may appear in ordinary cultures of some Zygomycetes when the general conditions are unfavourable to the fungus. For example, when Sporodinia grandis is transferred from its host to potato agar, it often happens that growth falls off after a few transfers; the number of matured reproductive structures then diminishes and both sporangia and zygospores may be replaced by abortive rudiments containing a yellowish pigment and the stock usually dies out. Phycomyces Blakesleeanus may show similar but less striking behaviour on unsuitable media. It seems a reasonable conclusion that the likeness between characters following irradiation and characters known to be indicative of unfavourable conditions for growth, shows that irradiation causes damage to the general balance of the fungi concerned.

We owe a very thorough study of the effect of X-rays on fungi to Dickson, who worked with both mycelia and spores. He records noteworthy results with *P. Blakesleeanus*, and with several species of

Chaetomium.

Six hundred subcultures from fifty-eight irradiated plates of *Phycomyces Blakesleeanus* produced seventeen sectoring colonies; eleven

^{*} Bean, W. J. and Brooks, F. T., New Phytol. XXXI (1932), 70.

[†] Dickson, H., Ann. Bot. xlvi (1932), 389. ‡ Nadson, G. A. and Philippov, G. S., C.R. Soc. Biol., Paris, xcm (1925), 473; J. Soc. Bot. Russe, xm (1928), 221. § Dickson, H., Ann. Bot. xlvi (1932), 389; xlvii (1933), 735.

variants were isolated from the sectors. Two, from heavily dosed cultures, did not form sporangia, and gave very few zygospores when mated with the appropriate strain of the parent form—this had not been irradiated. An orange pigment developed in the mycelium. Similar variants appeared in the progeny from less heavily dosed mycelia, so that dosage alone is not the decisive factor. Some variants from moderately dosed cultures were distinguished at first by the abundant and precocious crops of sporangia, but, after three months

of culture, these variants were reverting to the parent form.

More productive experiments were made with Chaetomium cochliodes. Preliminary observations on more than seven hundred cultures indicated that the stock had no marked tendency to vary in ordinary culture. Young and old mycelium was irradiated, and it was found that variations arose about two and a half times as often from old mycelium as from young mycelium, both having been equally exposed to treatment. Since subcultures from old mycelium, suitably irradiated, gave 92 per cent. variation, there can be little question of the efficacy of X-rays in bringing about change. Treatment of the mycelium of variants led to the appearance of still more variants, some of which seemed to have resumed normal characters. If true reversion did in fact occur, it is difficult to account for these results by supposing that the X-rays had knocked out a gene or destroyed a piece of a chromosome, but this matter cannot be discussed profitably in this place.

In all, Dickson observed some hundreds of variants, and thirty-eight were grown in pure culture; five of these were more or less unstable, the others seemed to be permanent. The changes affected the colour, dimensions, morphology, fertility and staining of the medium; some variants were more fertile than the stock, some less so, and some were sterile, but there was no general relation between increase of sterility and stronger staining of the medium. It appeared that the characters of the parent stocks were not changed in association with one another, except that, when fertility was much reduced, much whitish aerial mycelium might form; this phenomenon has already

been noted in fungi after treatment by heat.

Irradiation of suspensions of ascospores did not greatly affect the subsequent germination of the spores, but it had a pronounced effect on survival beyond the earliest stages, for with increase in the length of exposure to the rays there was marked decrease in the number of large mycelia formed. Many of the spores which did establish themselves yielded variants, and the number of variants increased with increase in the time of treatment.

There can be no doubt that X-rays provide a convenient and productive means of inducing variation in fungi. That this is so is well demonstrated by Dickson's investigation of seven species of Chaet-

omium; all species gave variants from treated spores, but all did not respond equally strongly. Two of the species were observed to form sectors in ordinary culture, but when the mycelia of these species were irradiated, one species varied seven times as frequently as the other. Dickson had already found that species of Fusarium, known to vary greatly in ordinary cultures, were not notably responsive to X-rays; this shows that instability under ordinary culture conditions is no indication that the fungus will react with special readiness to abnormal stimulation.

Dickson tested the effect of ultraviolet light on the mycelium and spores of *Chaetomium cochliodes*. The mycelium seemed to be unaffected, but treated ascospores gave about thirteen variant colonies for every hundred colonies subcultured. Many spores were killed outright. The variants were like those which appeared after the use of X-rays.

From many points of view the fungi are not the most suitable subjects for experiments on induced variation. Their simple organisation seems to be all in favour of reaction, and their rapid growth shortens the time necessary for experiments. On the other hand, the vegetative nuclei are small, and cytological work furnishes more conjectures than facts. Normal sexual processes are commonly absent, so that satisfactory breeding experiments are not possible, and though genetic investigations might be undertaken with some of the Phycomycetes, it is to be remembered that the resting spores of these fungi, formed after a sexual union, do not usually germinate readily. It is therefore fortunate that we can turn to work on induced variation in other organisms, and so supplement the impressions gained from work on fungi.

We may first consider the insects. In the latter years of last century some experiments were made by rearing larvae and pupae of Lepidoptera at high temperatures.* It was shown that by such methods some of the butterflies of Central Europe could be converted into forms comparable with races of the same species characteristic of parts of southern Europe. Heat treatment also yielded specimens with colour patterns suggesting a sort of generalisation of the patterns running through a number of species of the genus; it may be recalled that Dickson noted a tendency towards generalisation during his investigation of seven species of *Chaetomium*. There was some evidence that the induced characters in the butterflies were inheritable, but that side of the work was not pursued.

Much more recently, work on melanism in moths, by Harrison and Garrett,† has provided remarkable evidence on the induction of variation and its transmission to offspring. It has been known for a

^{*} Standfuss, M., N. Denkschr. schweiz. Ges. Naturwiss. xxxvI (1899), i, 1; Ann. Soc. ent. Fr. LXIX (1900), 82.
† Harrison, J. W. H. and Garrett, F. C., Proc. roy. Soc. B, XCIX (1926), 241.

long time that some moths which are light coloured in rural surroundings remote from towns, are represented in industrial areas by very dark forms, and the development of these dark forms appears to have gone on side by side with the growth of industrialism. Analysis of the smoke begrimed leaves of the food plants of some melanic moths showed that the dust contained lead and manganese. Light coloured moths were obtained from rural areas where melanism was unknown, and after breeding tests had shown that melanism was apparently absent from the stocks, larvae from these stocks were fed on contaminated foliage. Ultimately, melanic moths appeared in the insects so reared, and the melanism was found to act as a Mendelian recessive. This work provides clear evidence of the induction of variation and of the transmission of the variation through a normal

sexual process.

Flies belonging to the genus Drosophila have been much used in genetical work, and experiments have led to the induction of variation in members of this genus. Goldschmidt* procured a stock of D. melanogaster from Morgan, taken from a race whose history was well known. Mating flies were placed in flasks, and removed after eggs had been laid. Some lots of eggs were placed at 37° C. for some hours, and then kept at 25° C., to allow of further development; other lots were kept at a steady temperature of 25° C., a temperature very favourable to normal development. The results of these experiments showed many successes and many failures, in this respect agreeing with the results of high temperature experiments with fungi. The death rate was high, and of the flies which did survive, many were quite sterile, or only became fertile after a preliminary period of sterility; male sterility was specially prevalent. From the progeny of flies which hatched out, many variants were obtained, including specimens of some rare variants which had been seen only once before by any of the numerous workers on this well investigated insect. The variants included some non-transmissible modifications, and some which could transmit their characters; at times, flies hatched from heated eggs or larvae appeared to be normal, but abnormalities were found in their offspring, an interesting parallel with conditions known to occur in fungi.

H. J. Muller† and his associates have investigated the effects of X-rays on *Drosophila* very thoroughly, obtaining variants in large numbers and great variety. Induced sterility was of widespread occurrence, especially in males. It is well known that *Drosophila*

^{*} Goldschmidt, R., Biol. Zbl. XLIX (1929), 437.

† Muller, H. J., Proc. Nat. Acad. Sci., Wash., XIV (1928), 714; Genetics, XIII (1928), 279; Hereditas, XVI (1932), 160; Muller, H. J. and Altenburg, E., Proc. Soc. Exp. Biol., N.Y., XVII (1919), 10; Muller, H. J. and Mott-Smith, L. M., Proc. Nat. Acad. Sci., Wash., XVI (1930), 277.

varies spontaneously when grown in tubes with pieces of banana as food, but the suspicion arises that some at least of this supposed spontaneous variation may be induced by the crowded conditions, comparable with the methods used by the Chinese in dealing with goldfish. After treatment with X-rays however, the rate of change is increased enormously. Muller noted that the variants found most frequently in his work were like those which had appeared most frequently in the ordinary cultures of other investigators, this suggesting that some characters or groups of characters are more easily changed than others. One notable variant, distinguished by mottling of the eye, was unstable for that character, again furnishing a parallel with phenomena in some fungal variants.

The use of X-rays and high temperatures together further increased the rate of change, though it was clear that X-rays alone were much more effective than increases of temperature alone; there was however no evidence that the kinds of variants were in any way de-

termined by the agent used to provoke their appearance.

Tobacco plants have proved to be very responsive to X-rays, maybe because the cultivated plant is probably a hybrid, and therefore somewhat unsettled in constitution. A strain of tobacco which has been under observation for twenty-five years, and known to be stable in ordinary cultivation, has been investigated by Goodspeed* and his associates. Unopened flower buds were irradiated for about ten minutes and then allowed to develop; seeds from these buds gave rise to many abnormal plants, the variations affecting the stature of the plants, the shape of the leaves, and the colour, size and shape of the flowers. Some plants raised from seeds of treated flowers appeared to be normal, but, after selfing, variants developed in the progeny of these plants, still another example of delayed action. The plants often showed reduction in fertility, being at times quite sterile. Some plants were a mosaic of normal and variant tissue, and these plants may be compared with a sectoring mycelium. Cytological work showed that irradiation of tobacco plants may be followed by great disturbance in the behaviour of the chromosomes in dividing nuclei, and in the general construction of the nucleus.

Much interesting work on induced variation has been done on Antirrhinum. Baur started experiments in Germany as far back as 1908, and, in connection with this work, a stock of Antirrhinum has been carried on from year to year by selfing. Close observation of that stock has revealed little evidence of spontaneous variation, and it is of interest that the stock seems to have suffered no serious loss of vigour or diminution of fertility during the long period of inbreeding.

Among many experiments in which this stock has been used by * Goodspeed, T. H., Bot. Gaz. LXXXVII (1929), 563; Goodspeed, T. H. and Olson, A. R., Proc. Nat. Acad. Sci., Wash., XIV (1928), 66.

Baur* and his pupils, some experiments on seeds are of special interest. The seeds were soaked for twenty-four hours and then exposed to the emanations of radium. Little effect was produced unless the seeds were treated for at least forty-five minutes. With longer exposures, the seeds suffered loss of power of germination, and those which did germinate often produced short-lived seedlings with deformed cotyledons. Some plants which survived the seedling stage grew more slowly than normal plants, flowered late, and were specially liable to disease; others were abnormal when young, but seemed to become normal as they matured. A general relation appeared to exist between the duration of the exposure to radium and the extent to which the plants subsequently reacted, but the relation was by no means exact. Sterility was very common in plants grown from treated seeds, and even when abnormalities of morphology disappeared as the plants matured, sterility often persisted. Owing to this sterility, the plants were mostly propagated by vegetative means, giving clones which, commonly, retained the peculiarities of the variant from which the cuttings were taken. Anatomical and cytological investigations revealed a number of irregularities suggesting marked disturbance of ordinary development. The development of the pollen was sometimes so far affected that the process did not get beyond the earliest stages, and there were also indications that, after ovules had been laid down, they could then be replaced by ordinary vegetative tissue. However, some germinable seed was set by the variants, and this, on germination, usually gave normal plants of Antirrhinum.

Variants of similar character were obtained from another stock of Antirrhinum† after the material had been exposed to X-rays, ultraviolet light, high temperatures, and various chemicals. When young flowers were operated on, treatment was most effective if applied at a time when meiosis was probably in progress. None of the agents seemed to exert a specific effect. It is of note that temperatures of 47° C. soon killed active plant material saturated with water, for still another investigation; of Antirrhinum showed that a few pollen grains could function after they had been heated for three minutes at 116° C., provided that they were heated in a dry state after a thorough preliminary drying for several days in a desiccator. An even more surprising discovery was, that carefully dried pollen, after being heated at 86° C., for two days, was half as effective in bringing about fertilisation as was normal unheated pollen. As a further illustration of

^{*} Baur, E., Bibl. genet., Lpz., IV (1924); Z. Bot. XXIII (1930), 676; J. R. Hort. Soc. LVI (1931), 176; Stein, E., Z. indukt. Abstamm.- u. VererbLehre, XXIX (1922), 1; XLIII (1927), 1; Biol. Zbl., XLVII (1927), 705; L (1930), 129.

† Stubbe, H., Z. indukt. Abstamm.- u. VererbLehre, LVI (1930), 1, 202.

‡ Hiorth, G., Z. indukt. Abstamm.- u. VererbLehre, LVI (1930), 39.

the power of carefully dried plant material to resist heat, brief mention may be made of the work of Gain,* on the fruits of Helianthus annuus. A few of these retained sufficient vitality after fifteen minutes at 150°C. to begin to germinate, and flowering plants were raised from fruits which had been heated to 120° C.; these plants were unable to set seed.

It is however necessary to return to the experiments with Antirrhinum. 434 plants were grown from seed fertilised from strongly heated pollen; they included twenty variants, thirteen of these from pollen which had been heated to 100° C., or to higher temperatures. In contrast with these plants, 2771 plants from seeds fertilised by less strongly heated pollen, included only thirty-two variants. Of the fifty-two variants observed in these experiments, thirty-nine formed little or no pollen, and the plants had malformed anthers, various peculiarities of flower structure, and a tendency to develop very narrow leaves, a common

character in abnormal plants of Antirrhinum.

One further matter claims our attention, that of the influence of the age of reproductive bodies on the progeny developed from them. Gain (loc. cit.) found that similar abnormalities could be obtained in Helianthus annuus by strongly heating dry embryos, and by allowing embryos to dry and to age for long periods at ordinary temperatures. Still more recently it has been shown† that old fruits of Crepis tectorum may give abnormal plants; in one experiment, twenty-two out of twenty-seven plants from old fruits were abnormal. Cytological investigation of the root tips of these plants revealed many abnormalities in the nuclei, and further, the plants were shown to be chimaeras. The fruits used had been stored in the dry for from five to six years. It was then found that fruits of C. tectorum, ripened in the previous season and germinated after being heated for some weeks at about 55° C., gave many deformed plants resembling those grown from old fruits. Fruits of the previous season, after they had been heated in the dry for twenty days at 55° C., usually germinated as well as untreated material of the same age, but the seedlings often failed to get beyond the stage of spreading their cotyledons; of the plants that developed further, most were at first abnormal, but some ultimately assumed a normal appearance. Similar fruits gave a germination of over 70 per cent. after heating for forty days, but after forty-four days the germination was only 44 per cent. Seedlings from the more severely heated material never got beyond the cotyledon stage, and died after about a month. Fruits of the previous season, after treatment with X-rays, behaved much like heated fruits, and like old fruits after some years of dry storage; evidently, the diverse treatments were followed by similar response by the plant.

^{*} Gain, E., Rev. gén. Bot. xxxix (1927), 234, 306. † Navashin, M. and Shkvarnikov, P., Nature, Lond., cxxxii (1933), 482.

Cartledge and Blakeslee* have made comparable observations on a stock of *Datura* that had been under investigation for some years; this stock had a tendency to form aborted pollen grains, and, as that character was easily detected, it was used as a convenient means of estimating the rate at which the stock changed under special treatment. Old seeds, stored in the dry for from four to seven years, yielded many more plants with aborted pollen grains than seeds from one to five years old, similarly stored. There were indications that the tendency to produce abortive pollen was inheritable. The distribution on the plants of the flowers in which pollen abortion was or was not marked, indicated that most of the plants were composed partly of normal and partly of altered tissue; that is, the plants were comparable with a sectoring mycelium; few plants consisted wholly of altered tissue.

In Helianthus, in Grepis and in Datura, it seems that, during storage, or under the influence of special treatment, changes occurred in at least some of the cells of the embryo, and that, at any rate in Grepis and Datura, these changes resulted in the development of plants of heterogeneous composition; this suggestion however still awaits com-

plete demonstration.

Abnormalities in seedlings grown from aged seed bring us back again to the fungi, for changes in pigmentation of the conidia have been recorded† in species of Aspergillus grown from old spores. Unusually blue conidia developed on colonies of A. versicolar started from conidia two years old, and conidia nine years old, taken from an apparently normal strain of A. glaucus, produced colonies with colourless conidia. The nine-year-old spores did not germinate well, and here, presumably, we have an example of variants derived from moribund spores, a circumstance reminiscent of some of the results obtained in experiments with high temperatures.

The foregoing survey is by no means complete, but it is believed to contain a fair selection of the facts at present available. The selection of facts has been made in order to show the kind of evidence that is available, not to support any special point of view, except the underlying idea that it is possible to induce variation by experimental means. Reference has not been made to the extensive literature on induced variation in bacteria and in many of the lower animals, and no attempt has been made to discuss the large volume of work on the extensive variation shown by some fungi under apparently ordinary conditions of culture. It is however by no means impossible that some of this so-called spontaneous variation may be traced ultimately to

^{*} Cartledge, J. L. and Blakeslee, A. F., Proc. Nat. Acad. Sci., Wash., xx (1934), 103.
† Blochwitz, A., Ber. dtsch. bot. Ges. x11 (1923), 205.

reactions following on the wounding caused by using pieces of hyphae to start new cultures, and some may be due to the use of media which are really unsuitable to the needs of the fungus. It is usual to assume that media which serve the needs of a particular investigation in providing desired stages in the life-history of a fungus are favourable media for the growth of the fungus but it does not follow that the needs of fungus and investigator necessarily coincide. It is well known that many fungi are stable in culture on such media as potato agar and unstable on synthetic media; the latter, convenient as they are, probably lack some constituent necessary to healthy growth, or contain some substance or combination of substances which, while promoting strong growth, also upset the general balance of the developing organism.

It is natural enough to seek for an explanation of the facts, but it is doubtful if we are yet in a position to begin to suggest one; as the phenomena are far from simple, it is probable that they cannot all be brought under one explanation. Yet, as a conclusion to this provisional survey of a wide field, a few general remarks appear to be

necessary; they need not be extensive.

There can be no reasonable doubt that stocks of organisms, known to be stable under ordinary treatment, yield, after certain kinds of experiments, forms which have not been noted before in the stock. Since these forms appear only after experimental treatment, it is not easy to avoid the conclusion that they owe their origin to some effect of the treatment. It has been suggested, and there seems to be no decisive argument to negative the suggestion, that the variants are nothing more than forms of the stock, rarely seen under normal treatment, but selected by the conditions of experiment. It is difficult to accept this view, at any rate for the fungi. Some of the variants differ so conspicuously from normal that they could not be overlooked, did they occur at all frequently in ordinary cultures. The spores of some variants germinate as readily as the ordinary spores, they need no special treatment to cause them to germinate, and the variant colonies do not demand special treatment for their healthy growth. In mixed culture, normals and variants grow side by side and retain their characters, so that they can be distinguished with the utmost ease even in dry cultures which are months or years old. Such variants could hardly escape observation over a period of years if they were normal, though occasional products of a normal strain of fungus. The idea that the variants are selected by the treatment becomes almost absurd when it is recalled how freely variants appeared in Chaetomium after treatment with X-rays.

Certain features come out again and again in the results of work on induced variation, and, so far as the peculiarities of their organisation allow, the fungi fit well into the general scheme. Two outstanding features are, a common but by no means universal reduction in fertility, and a frequent but not invariable weakness in growth; such features suggest that the variants are damaged versions of the normal stocks from which they have sprung. This interpretation is well supported by the abundant evidence that the induction of a variant is often followed by a period of adjustment, a period not always ending in the same way. Sometimes the adjustment cannot be made, and the organism dies, maybe as a sporeling, maybe after it has passed through several transfers: the damage has been so severe that it cannot be repaired. Sometimes adjustment is so complete that the normal form is regained, either in the first culture or after a period of growth: the disturbance has been transitory. Sometimes there is an intermediate condition, and adjustment leads to the establishment of a permanent variant: the damage has caused a shift in the general

balance within the organism.

The nature of the change remains obscure. Much time could be spent in speculations about gene changes and alterations in the chromosomes. Such changes may, and probably do occur. It seems highly probable that some nuclear change, and some mixing of nuclei of different qualities must be concerned in a sectoring mycelium. and in induced chimaeras in higher plants. But, until we have more definite evidence it is unwise to attribute everything to nuclear changes; the similar effects which follow heat treatment and age for example, suggest that a general derangement of the physiological balance of the cell may well be responsible. The likeness that seems to exist between the effects on cells of violent external treatment and of the slow changes which must proceed in resting spores and seeds, is perhaps the most interesting feature of work on induced variation: the violent external agents may well hasten the normal changes of a degenerative nature which presumably assist in bringing about the death of a spore or seed unable for some reason to germinate. We have at least a glimpse of a possibility that some new forms may arise from aged material and establish themselves. If this be so, the evolutionary process may depend in part on the running down of the biological machine.

COOKE'S ILLUSTRATIONS OF BRITISH FUNGI

The eight volumes of Cooke's *Illustrations of British Fungi* are much prized, but referred to with increasing doubt as a wider knowledge of agaric species is obtained. The fact is that a large proportion of the plates are wrongly or doubtfully named. For the most part the figures are well done; there are of course some poor figures, but the whole forms a very valuable series, perhaps the finest set of illustrations of agarics in existence. Many modern reproductions are superior, but

are scattered in various publications not easily acquired.

In the Transactions of this Society there have been two lists of criticisms of the Cooke plates. C. B. Plowright (Trans. Brit. mycol. Soc. I (1898), 39-40) made a few comments in his presidential address, but suggested only a few changes in the names given by Cooke. Emile Boudier (II (1906), 150-7) made some rectifications and observations which were very illuminating. Cooke (III (1907), 26-9) replied, agreeing to some of the determinations but vigorously disputing others. Since then there have been numerous citations that sometimes have referred to species other than what Cooke named them, but there has been no paper covering all the plates.

By a fortunate chance my friend, M. Joachim, a past President of the Société Mycologique de France, showed me the copy of a manuscript which had been written by Lucien Quélet. It bore the title "Annotations d'après le Dr Quélet des planches de l'ouvrage: Handbook of British Fungi by M. C. Cooke, second and revised edition. London 1883". The title is erroneous, as the annotations were not of Cooke's Handbook, but of the *Illustrations of British Fungi* (1881–91). The manuscript contains Quélet's views of all the plates. Where he makes no comment, it is to be presumed that he agrees with the name

printed on the plate.

The manuscript appeared to be of so much interest that I asked M. Joachim for permission to have it reproduced in these *Transactions*.

It then occurred to me that the work could be made much more useful if supplemented with a review of Cooke's *Illustrations* by modern mycologists well qualified to undertake this task, and I therefore referred to the two eminent authorities, Prof. Dr René Maire and Mr Carleton Rea. They both very kindly agreed to annotate the plates, and the result is here given in tabular form. In frequent cases there will be found a conflict of opinion. This was inevitable. Even with the living fungus, eminent mycologists do not always agree. How much more must this occur with an illustration which may be an imperfect representation of the original? The true identity of many of the agarics figured by Cooke can never be known with certainty.

M S

Dr Maire, in his manuscript, used many of the new generic names which are now generally adopted by French mycologists. These, however, are not printed, as the purpose of the annotations is sufficiently served by using the specific epithets only, except where clearness requires the genus to be shown.

In the columns under both Maire and Rea, a dash does not indicate agreement as in the Quélet column, but means that no opinion

is ventured.

The annotations could have been made more complete by including the many citations from monographs, etc., that have appeared in recent years. Doubtless this would have added interest and would have revealed further differences of opinion. I doubt, however, whether it would have added to the practical value of the paper. All will continue to have their own views about some of the plates, but the new light that is now thrown upon Cooke's *Illustrations of British Fungi* will be welcomed by students of the agaries everywhere.

The species have been listed in the rotation of the bound volumes as finally issued. This does not correspond with the numbering on the plates. There are many volumes scattered about the world which have been bound differently, so an index to the plate numbers has been added which will give further value to the present paper.

A. A. PEARSON.



COOKE'S ILLUSTRATIONS OF BRITISH FUNGI

No. of bound volume	No. printed on Plate		Cod	OKE	Quéler
. 1	I 2	Agaricus	(Amanita)	virosus phalloides	solitaria on junquillea
3	3	Additional	Specialists	vernus maþþa	ou citrina var. albata
4 5 6	117 6	anners.	Section 2	muscarius pantherinus	2
7 8 9	7 8 277	-	Ministrative Mi	excelsus strobiliformis strobiliformis	virescens (excepté le bulbe) non! incontru—rubens (?) ou solitaria
11	69 69	EFFICE.	Marina	rubescens spissus	2
12	70 10	and the same of th	generally distances	nitidus asper	aspera var. pallescens
14 15 16	34 11 12	-	Miles of	magnificus megalodactylus vaginatus	Lepiota guttata (?)
17	13 35	- Management - Man	-	strangulatus adnatus	malé (?) junquillen vetustior ou aspera pallescens
19 20 21	21 22		(Lepiota)	procerus rhacodes	trop petitmastoidea
22	23 28	anada	Service Servic	excoriatus gracilentus	Microsel Hallongel
23	24		State Address	mastoideus	représente mieux cristata
24 25	14 25	-	Services Services	acutesquamosus Badhami	erhalics
26 27	26 37	*******	property	meleagris biornatus	nangka manag
28 29	27 38		******	hispidus	non!
30	39	hanne	- Annual - A	clypeolarius metulaespoyus	? clypeoloria, gracilis avec ét maladif des lamelles
31 32 33	29 40 36	Mariness Mariness	Minneys Minneys	cristatus ermineus Vittadini	3
34 35 36	41 15	-		holosericeus naucinus	ou plutôt guttata (?)
	5	, 	-	cepaestipes	lutea et cepaestipes
37 38 39	42 43 18	Totales		carcharias cinnabarinus granulosus	Asseque Spranse Onstager
40	213	_		granulosus var, rufescens amianthinus	Manue

forme de spissa, teinte trop verte peut-être A. Emilii Riel solitaria rubescens spissa mais atypique. La figure du haut rappelle plutôt aspera (?) mauvaise figure de echinocephala aspera (forme à verrues grisâtres) anomalie de rubescens lenticularis var. eguttata vaginata inaurata Amanita gemmata Fr. (=junquillea Q.) procera, petite forme rhacodes excoriata gracilenta? trop petit et trop mince, peut-être L. cepaestipes var. nigricans Bagl.	ypical verna (syn. virosa) halloides, with typical free-lobed volva halpa var. alba Gill. halpa (syn. citrina) huscaria hustaria. It has the striate margin and con- centric rings to apex of volva kcelsa hobiliformis typical hubescens hissa poor hubescens poor hubescens poor hubescens poor halloid (syn. inaurata) hada (syn. junquillea and hada (syn. junquillea and hacodes hacodes hacodes hacodes hoor gracilenta, does not show minute scales on	3 4 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22	3 4 117 6 7 8 277 9 69 70 10 34 11 12 13 35
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L. cepaestipes var. nigricans Bagl. acutesquamosa Badhami (spores, données d'après E		1	28
L. cepaestipes var. nigricans Bagl. acutesquamosa Badhami (spores, données d'après E	stem	1	
Badhami (spores, données d'après E	ot mastoidea, does not show excoriated margin or scales on stem	23	24
Badhami (spores, données d'après E	cutesquamosa, large form	24	14
massec, hop petites)	Padhami typical	25	25
	eleagris	26	26
très voisin d'un <i>Lepiota</i> fréquent à b Alger, mais dont la chair ne rou- git pas	iornatus	27	37
	ypeolaria type	28	27
	ypeolariodes Rea	29	38
	ypeolaria	30	39
	ristata	31	29
	minea	32	40
* // /	ittadinii, but should give gills finally greenish	33	36
	aucina, hot bed form	34	41
	tiea	35 36	15
	epaestipes	30	5
	ircharias	37	42
	innabarina	38	1
	ranulosa	39	18
	archarias ranulosa var. rufescens	40	213

No. of bound volume	No. printed on Plate	Co	OKE	Quélet
41	30	Agaricus (Lepiota)	polystictus	Armill. rufa, gracilis? Arm. luteo-vireus, gracilis?
42	85	gan rapa santripi	sistratus	Africa serv
43	19	specialists unitable unitable unitable unitable unitable unitable unitable	mesomorphus seminudus Bucknalli	et var. lilacina Quél.
44 45	44 118	parties and displayed and disp	medullatus gliodermus delicatus	copié
46 47	17	Maringon Shrandari Maringon Shrandari	lenticularis Georginae	haematosperma, forme très grèle
48 49 50 51 52 53 54	20 245 31 33 86 71 45	— (Armillaria)	bulbiger focalis — var. Goliath aurantius robustus var. minor ramentaceus haematites	? subannulata Batsch carcharias, forme malade
55 56 57	46 32 47		constrictus melleus subcavus	holosericea cepaestipes ou serena
58 59 60	16 72 53	_ (Tricholoma	mucidus) equestris sejunctus	malè
61 62 63 64	54 73 74 55		porteniosus fucatus quinquepartitus resplendens	Coll, grammocephala Buil.
65 66 67	8 ₇ 75 76	Manager and American Windows and American Stronger American	spermaticus colossus acerbus	columbetta columbetta, vetusta malè
68 69 70 71 72 73 74 75 76 77 78 80 81	56 57 58 197 88 198 214 59 48 215 199 60 61		nictitans fulvellus flavo-brunneus albo-brunneus ustalis stans rutilans luridus guttatus columbetta scalpturatus imbricatus imbricatus immundus	fulvum fulvum ou ustale grêle malè striatum ? rappelle Coll. maculata luridum
82	49	-	murinaceus	mention
83 84 85 86	50 165 51 90		terreus — var. argyraceus atrosquamus ori-rubens	hordum orirubens ou argyraceum malè

Maire	REA	No. of bound volume	No. printed on Plate	
?	polysticta	41	30	
*****	sistrata	42	85	
seminuda Bucknallii, mauvaise figure	seminuda I doubt Bucknallii though I know it well	43	19	
glioderma	lookslike Fries Iconesfigure glioderma	44 45	44 118	
lenticularis Georginae	lenticularis Georginae	46 47	17 132	
bulbiger — robusta ramentacea	bulbigera — robusta ramentacea	48 49 50 51 52 53	20 245 31 33 86 71	
haematites constricta	like the original painting of haematites	54	45	
mellea Lepiota Brebissoni Godey ou Magnusiana P. Henn., Maire	constricta typical mellea —	55 56 57	46 32 47	
mucida trop pâle Tr. sejunctum, mauvaise figure	mucida equestre possibly sejunctum but not typical	58 59 60	16 72 53	
Coll. platyphylla ? ? ?	plaiyphylla	61 62 63 64	54 73 74 55	
? Tr. colossus suffocatum	spermaticum, sp. verrucose acerbum, but too deep in	65 66 67	8 ₇ 75 76	
flavobrunneum ?	colour — —	68 69	56 57	
pessundatum, mauvaise figure albobrunneum, mauvaise figure ustale albobrunneum? malé rutilans	possibly pessundatum albobrunneum typical ustale stans rutilans	70 71 72 73 74	58 197 88 198 198	
? orirubens? ou forme voisine columbetta? atrosquamosum, mauvaise figure imbricatum	luridum poor guttatum (Schaeff.) Rea columbetta certainly not scalpturatum imbricatum poor	75 76 77 78	214 59 48 215	
vaccinum T. immundum=T. fumosum Pers. non Fr.	vaccinum typical Collybia fumosa Pers.	79 80 81	60 61	
murinaceum Fr. non Quél., mauvaise figure	murinaceum typical	82	49	
? scalpturatum= argyraceum atrosquamosum ? orirubens	a form of terreum argyraceum typical terreum var. atrosquamosum orirubens (? diseased form of terreum)	83 84 85 86	50 165 51 90	

No. of bound volume	No. printed on Plate		C	OOKE	Qualet
87	278	Agaricus	(Tricho	loma) macrorhizus	*** *** *** *** *** *** *** *** *** **
88	91			saponaceus	malè
89	216	-	Antonia	id. var. stipite-	V man
				squamuloso	5
90	166	Mana	personal and the same of	cartilagineus	triste
91	52	-	domina	atrocinereus	Conve
		-	**************************************	cuneifolius	
92	261	******	100000-004	cuneifolius var.	? forme de Russula lilace
93	92	-	-	cinereo-rimosus	1 21 - 11
94	93	-	*****	crassifolius tumidus	Georgii
95	167	-		virgatus	hordum?
96	62	-mran	********	sulphureus	who we
97	181	arriving.	***************************************	bufonius	y whitens
98	94		Reference .	lascivus	
99	217	#F-1414mg	Finning	id. var. robustus	album, gracile?
				Rt. vai . fobusius	Omphalia gilva
100	77	-	******	inamaenus	copié de Fries
101	95	-	Province	ionides	malè
102	06	Mark year	***************************************	cerinus	: Marasmitis Oreades
102	96		Whiteeda	carneus	mal copiés
103	63		-	coelatus [
103	262		Per Strang	gambosus	Clitacybe gentropa
105	229	_	*******	amethystinus	non! luridum ?
106	64		-	albellus	Georgii, malè
107	168	-	-	tigrinus Colomondoni	non! copié de Fries
108	279	*********	No.	Schumacheri patulus	forme de nebularis
109	218	-	piones.	arcuatus	Sharangi
			anning.	oreinus	A manual Lit
110	65	WHEN AND THE PERSON NAMED IN	Married	albus	? male
III	78	-	- Parentage	leucocephalus	non! Cort. sebaceus?
112	169	-	Web organic	militaris	quelque chose de
	CC				acerbum
113	66	attending.	. The strange	personatus	malé
114	67	-	*******	nudus	malé. ?
115	133		**********	id. var. major	malè
116	170	Managed	Minney	cinerascens	malè
117	97	-	- Military	panaeolus	ASSESSO
118	98		Minimum	grammobodius	NAME OF THE PARTY
119	119	-	Printeg	melaleucus et var.	-W-Mer
120	68			porphyroleucus	
121	1	Manage .	-	brevipes	?
	99	-	War way	humilis	humile?
122	263			7	sordidum
1	3		-	humilis	malè
123	171			exscissus	
124	219	Million		subpulverulentus	anneau.
125	100	******	*****	sordidus	number 1
126	120	-	who the same of	paedidus	malè
		-	Ministra .	lixivius	car malala
127	172	-	***************************************	putidus	ou melaleucum malè
	79 80	_ (Cl	itocybe)	nebularis	mate
129			Tribus.	clavipes	No.
130	246	-	-	inornatus	Territoria.
191	264	-	-	hirnaeolus	? leucophylla
31	204	Printer.	-	cyanophaeus	forme de sordidum
34	134		-	amarus	we will the surging
	1		-	socialis	flaccida

Maire	Maire Rea		
?		87	278
saponaceum, mauvaise figure	saponaceum	88	91
saponaceum var.	id. var. stipite-squamuloso	89	216
terreum	a form of terreum	90	166
atrocinereum cuneifolium	atrocinereum cuneifolium	91	52
cuneifolium forma	var. cinereo-rimosum	92	261
Georgii forma?	not crassifolium of my work	93	92
virgatum	virgatum	94	93 167
sulphureum	sulphureum	95 96	62
id. var. bufonium	bufonium	97	181
? trop grêle		98	94
Clitocybe Alexandri Fr. = C. gilva Quél. non Fr.		99	217
inamaenum	inamaenum	100	77
ionides	ionides	101	95
*	not cerinus		50
carneum	carneum typical	102	96
caelatum	caelatum typical		
?	gambosum	103	63
		104	262
Georgii, trop jaune	-	105	229
inspiré de Fries		106	64 168
Clitocybe nebularis	not patulum	107	
forme de brevipes	brevipes	109	279 218
graminicola (Velen.)	- Ortotpes	109	-10
beaucoup trop jaune. Clitocybe?	album, extreme form	110	65
	leucocephalum	III	78
	militare	112	169
saevum (= personatum var. anserinum Fr.)	personatum type	113	66
do.	saevum	114	67
do.	personatum	115	133
?	cinerascens poor	116	170
Panaeolus	panaeolus	117	97
grammopodium	turritum	118	98
melaleucum	melaleucum	119	119
7	id. var. porphyroleucum		60
brevipes	brevipes	120	68
melaleucum var. phaeopodium	sordidum	121	99
sordidum, mais lamelles trop pâles ? ?	humile	122	263
melaleucum var. excissum, pâle	melaleucum	122	203
melaleucum var. excissum?	excissum poor	123	171
	subpulverulentum	124	219
sordidum, mauvaise figure	sordidum	125	100
	_	126	120
	_		
?		127	172
nebularis	nebularis	128	79 80
clavipes	clavipes poor	129	
inornata	inornata	130	246
	1	1	
Panus torulosus jeune?		131	264

No. of bound volume	No. printed on Plate	(looke	Quélet
133	265	Agaricus (Clitoc)	be) venustissimus	copiés
134	101	phosphalia phosphalia	odorus	S Sidney Wilde
135 136	102	Mineral pro-copy	trogii	forme de viridis décoloré
136	200	Married Married	rivulosus	ventosa
	1	-	id. var. neptuneus	absolument faux! ? pyxida
137 138	121	Minutes - Process	cerussatus	notes a second of the second o
138	122	frenies temp	id. var difformis	Medicals
139	18	99000	phyllophilus	agen p
140	103		pithyophilus	Melinia
	0	*****	tornatus	Minimal Marie Mari
141	82	\$10.00A	candicans	phyllophila
142	104	-	dealbatus	cerussata
143	173	and see	id. var. minor	Material
144	174 182	-	gallinaceus	phyllophila
145 146	280		aggregatus	No. openia
140		*******	elixus	gilva Pers.?
147 148	175	Manage of State of St	fumosus	divines.
149	105		tumulosus	apon.
150	106	Ministra Ministra	opacus	?
151	135	Mileson Mileso	giganteus	mbr/glockly
152	107		maximus	geotropa
153	281	Maried Maried	infundibuliformis incilis	St- street
	7 -1	-	1/1611172	?
		-	parilis	
154	83	derina derina	geotropus	trop jaune
155	177	Service Service	id. var. subinvolutus	with drawn
155 156	177	derman decents Notices employe minimum decents decents decents decents decents decents decents decents	subinvolutus	property and the second
157	136		gilvus	geotropa
0				
158	109	-	splendens	Sin May
159	84	generally braining	flaccidus	MICHAL END
100	123	Million British	flaccidus	Attinop
161	107			
162	137		lobatus	Patricy
163	111	Worked Worked	senilis	Allehydag
164	112	- Company	catinus	Nedtron
165	138	-	tuba	"Miljanuda
165 166	113	Ventural Martinal Mar	ericetorum	Widnes .
			cyathiformis	Phylin _{tic}
167	220	Marine Marine	expallens	
168	230	- Minney America	obbatus	cyathiformis
169	231	Perhaps Projects	pruinosus	expallens?
			In antivaria	squamulosa
70	114	Marine Marine	brumalis	Prilyens
71	115		metachrous	
72	115	-	ditopus	Notice
73	232	Arrivan	diatretus	Marian.
74	124		fragrans	?
75 76	125		angustissimus	Without
76	² 33		obsoletus	J. 10 .
77 78		-	ectypus	dealbata
78	183		bellus	and the same of th
79 80	139	- (<u> </u>	laccatus	Callat:
81	127	-	Sadleri	Collybia
82	140	- (Collybia)	radicatus	- Managaria
83	201		longipes	- X - X -
~o	128		blatablatter	× 7

Maire	Rea	No. of bound volume	No. printed on Plate	
Pleurotus olearius?		133	265	
venustissima	venustissima			
odora, mauvaise figure	odora poor	134	101	
odora var. Trogii	viridis	135	102	
rivulosa?, mauvaise figure	not rivulosa	136	200	
Omphalia pyxidata	-	-3-		
cerussata	cerussata	137	121	
to come the page		138	122	
phyllophila	phyllophila	139	81	
gallinacea	- projetopitita	140	103	
tornata	tornata	140	103	
gallinacea?	phyllophila	T 4 T	82	
		141		
ressemble un peu à connata	cerussata not typical	142	104	
T have	dealbata var. minor	143	173	
gallinacea	gallinacea	144	174	
aggregata	aggregata	145	182	
inornata	inornata	146	280	
Col. fumosa Fr. non Quél. nec Bres.	Tr. cinerascens	147	175	
tumulosa	tumulosa	148	105	
Infrare		149	176	
gigantea	gigantea	150	106	
geotropa forma maxima	maxima	151	135	
infundibuliformis	infundibuliformis	152	107	
Manage	incilis poor, margin should	153	281	
	be crenate	-		
parilis	parilis typical			
geotropa	geotropa	154	83	
geotropa, mais spores inexactes	subinvoluta	155	177	
geotropa		156	177	
infundibuliformis forma gibba Fr. Mon. p. 119	infundibuliformis	157	136	
splendens	splendens small form	158	109	
inversa	inversa	159	84	
inversa (C. flaccida, insuffisamment distinct d'inversa)	flaccida	160	123	
inversa	lobata	161	137	
2	viridis	162	110	
catinus, trop blanc	catinus	163	111	
tuba	tuba	164	112	
ericetorum? ou Hygrophorus niveus?	ericetorum	165	138	
cyathiformis	cyathiformis, but does not	166	113	
	show reticulate stem		- 3	
cyathiformis, forme grêle?	expallens	167	220	
5		168	230	
?	cyathiformis var., stem is right	169	231	
brumalis	brumalis, but does not show the finally yellowish gill	170	114	
metachroa	metachroa small form	171	115	
ditopa	ditopa	172	116	
diatreta	diatreta	173	232	
fragrans	fragrans	174	124	
angustissima	angustissima	175	125	
2		176	233	
cyathiformis	cyathiformis	177	126	
c yashing of nites	- Cyanagornas	178	183	
Tanamia laggata at war amathusting	laccata and var. amethystina	1		
Laccaria laccata et var. amethystina		179	139	
fasciculare, forme stérile	fasciculare		127	
radicata	radicata	181	140	
longipes	longipes	182	201	
platyphylla	platyphylla	183	1. 138	

No. of bound volume	No. printed on Plate		C	COOKE	Quilet
184	292	Agaric	us (Collyb.	ia) semitalis	made a service and a service as a service as
185 186	141	a	-	fusibes	and the second s
	142	********	Man cin	maculatus	CHARLES
187	221	was before	V	var. immaculatus	
188	282	No.	* Amening	distortus	
189	143	Bulletone .	Second	butyraceus	and the second
190	202	Boothing	*****	xylophilus	make and
191	184	· per-prise	Militaria	velutipes	W. DV 2 ·
			- Maria N	laxipes	
192	129	E PPorts	Youver	mimicus	1
193	149	Service -	XXX	vertirugis	dryophila
00	13	Personal	All Street Street	stipitarius	Amenda
194	150	Magneton	eren.		
-34	-30	- Britana	Marries .	hariolorum	Name of pr
105	283	Atomie	Married Marrie	confluens	
195	130			ingratus	
		Pitroin.	Miles Park Spa	conigenus	****
197	144		Mr. on	tuberosus	No. of the Control of
		object and	Arturaryi	cirrhatus	
198	205	almosts.	#FFFFFF	collinus	malè
199	145	incomp.	Service and	ventricosus	111/11()
		Philips.	Military.	Stevensoni	
200	266	Anna	950.00	psathyroïdes	
201	203	Phone	Her up	xanthopus	forme de Mycena micea
202	146	Windows	Phone		dryophila
203	151	to word	******	nitellinus	extuberans
7-3	-3-	-	Phone .	succineus	non?
204	152	-		nummularius	aquasa
204	134	******	Men for our	esculentus	14 404
205	267	Person	- AMONG	tenacellus	(Comp.
205		Princip	beinging	acervatus	malė
	204	Process	Protes	dryophilus	language and the second
207	234 268	Accessed	Person	aquosus	? deyophila
208	208	Process	199.0146	exsculptus	Toronto.
		Minne	Window.	macilentus	dryophila
209	147	-	Processings	clavus	
		Manager .	Marina	ocellatus	Mycena ucuularis
		Photograp	Pr. No. ton	muscigenus	Mycena flavo-alha
210	153	-		rancidus	Mycena lactea
		Methodological	deletera.	coracinus	***.ue
211	154	Pitropa.	Marining.	inolens	944
	7	-	Windowski	plexipes	Pi-15 to
				prexipes	rancida
212	155	-	April Salas	atratus	
		-	At Your	ambustus	wine da.
213	269	-	two-mag		defense
214	270	-	Witness	laceratus	Hygrophorus distortus
- 1		-	infrage.	protractus	gending
215	247	-		tesquorum	- Million and American
	-1/		Montres	tylicolor	Adding
		-	-	clusilis	
216	156	-	(Mycena)	pelianthinus	Warries
		-	(balaninus	malè
217	284		Marrie -		PARTIES.
		-		elegans	Shranga
218	131		-	rubro-marginatus	sena, r
		-		strobilinus	coccinea
	1	-	phone	- var. coccineus	2000cgs
219	157		-	roselius	PATES.
220	158		attended.	purus	Williams
	-30	-	******	pseudopurus	pura
		-		zephirus	F

Maire	Rea	No. of bound volume	No. printed on Plate	
semitalis	semitalis	184	292	
fusipes	fusipes	185	141	
maculata	maculata	186	142	
?	maculata var. immaculata	187	221	
distorta	distorta	188	282	
butyracea	butyracea	189	143	
7		190	202	
velutipes	velutipes	191	184	
3	laxipes	191	104	
	mimica	192	700	
Marasmius undatus	undatus	1	129	
		193	149	
Crinipellis stipitarius	stipitarius	1		
	both hariolorum	194	150	
confluens	CONTRACTOR OF THE PARTY OF THE	1		
acervata vieux	acervata	195	283	
conigena	conigena	196	130	
tuberosa	both tuberosa	197	144	
cirrhata Cke. non Schum. = C. cirrhata var. Cookei Bres.		-57		
description .		198	205	
radicata forma	persona	199	145	
	-	- 33	-43	
Mycena sp.	<u> </u>	200	266	
dryophila	the have			
	xanthopus	201	203	
extuberans? certe non nitellina	extuberans	202	146	
i' dryophila forma aquosa	succinea rather too dark nummularius	203	151	
clavus (sensu Quélet)	esculentus esculentus	204	152	
clavus (sensu Quélet)			-6-	
acervata	acervata	205	267	
dryophila	dryophila	206	204	
dryophila forma aquosa	dryophila var. aquosa	207	234 268	
dryophila var. funicularis		208	268	
dryophila var. funicularis	dryophila var. funicularis	1		
Mycena acicula	Mycena clavus	209	147	
Collybia cirrhata	ocellata			
Mycena lactea	muscigena		1	
rancida	rancida	210	153	
coracina	coracina	710	-33	
inolens? rancida?? (lui ressemble mais dit inodore)	inolens plexipes	211	154	
atrata	atrata	212	155	
ambusta	ambusta	1	-33	
lacerata?	not lacerata	010	269	
tacerata:	1	213		
N/MANAGE	protracta	214	270	
er-nodes	tesquorum			
	tylicolor, the stem is white pruinose	215	247	
clusilis	clusilis typical			
pelianthina?	pelianthina	216	156	
Marasmius cohaerens	balanina? = atro-marginata			
Langei?	?	217	284	
rubromarginata	rubromarginata	1	1	
strobilina	strobilina	218	131	
coccinea	coccinea	7.7		
rosella	rosella			
	pura	010	157	
pura	pura	219	157	
pura forma				

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221	185	Agaricus	(Mycena)		floridula
		Monte	tra	lineatus	(Frage)
222	159		No-same.	luteo-albus	PE-14
		w/water	bradess	flavo-albus	(Avi en
		Moderate	Almone	lacteus	Acuto
223	235	*****	ermin di	proliferus	trop jaune
224	148 186	mortuus.	Wildelia	excisus	de hope
225	100	amount	Market of	psammicola	filopes on debilis?
226	206	-	******	rugosus	ammoniaca
227	222	-		sudorus	galericulata f. alba
22/	222		ment and	galericulatus — terrestrial var.	of track also. May define
228	223	Menon	personal, 1	var. calopus	Mighton Milyon in
~~0	~~3	****	Manager .	polygrammus	Million in
229	224	- Manager Mr	******	parabolicus	e of
3	774	Anthors	-	tintinabulum	eNeerhi
230	285	Service .		dissiliens	El Ante
	3	pilos y P	months	plicosus	direction and a second
231	236	*****	desired rel	pauperculus	cyanorhiza?
J	-3-	. Bretone	N/ALVAN	atro-cyaneus	Cyanoraeza;
232	237	-	-	pullatus	galopus
233	237 187	-	to the same	leptocephalus	gatopus
00	,	Minus.	Withing	alcalinus	ammoniaca
234	225	*****	***************************************	alcalinus	ATTETER/FERRECES
235	238	Monthe	*****	ammoniacus	Wroten
	00	*****	Property	metatus	WAY W
236	188		Newsloom -	aetites	malė, trop Jaune
		-	Green pajac Direktoran Green and	stanneus	collariata "
237	160		-	vitreus	? malè
000	161		-	tenuis	Militaria
238	101	-	*****	filopes	Wholes
000	286	Personal	******	Iris	Specifies.
239			(Administry)	amictus	*servides
240	189		~~~	debilis	lutenalba
		-		vitilis	min-mi
		-	(MARGAN) (MARGAN) (MARGAN)	collariatus	non. lineata?
241	190	-	Name .	speireus	malè, pas jaune
-1 10		Photosop	*****	tenellus	Omphalia stellata?
040		No. of Street,	******	acicula	malė, pas rosė
242	162	Time to	person, p	haematopus	Televisia.
243	163	-	Manage .	cruentus	derinos.
-43	103		-	sanguinolentus	%SMage
244	207			crocatus	Working
-11	20/	-		chelidonius	entencia America America America
245	208	-	Arriva	galopus	- Martine Mart
13	-00		-	epipterygius	drawaya
246	191		-	clavicularis	Person
. *	-3-	-		pelliculosus	Provide N. d.
247	248	-		vulgaris	rorida
	7.	-	1	citrinellus blicato	WARRY
		-		plicato-crenatus	Minimum
248	249	-		roridus	emptype.
	TJ	-		stylobates	Medical
				tenerrimus electicus	Militaria.
249	192			sacharifan	
77.				sacchariferus	Manage
	1	-	-	discopus	

Maire	Rea	No. of bound volume	No. printed on Plate
Adonis	Adonis	221	185
ineata lavo-alba ? lavo-alba ?	lineata luteo-alba flavo-alba	222	159
actea var. pitya	lactea prolifera	223	235
galericulata vieux? debilis?	Berkeleyi, but unknown to me	224 225	235 148 186
rugosa galericulata forma alba galericulata	rugosa sudora (viscid on pileus) galericulata	226 227	206 222
? inclinata (sensu Quélet) bolygramma	galericulata galericulata var. calopus polygramma	228	223
		229	224
	dissiliens plicosa	230	285
? atro-cyanea	atrocyanea	231	236
alcalina?	pullata	232 233	237 187
ammoniaca inclinata en haut inclinata en basouforme de galericulata	inclinata typical alcalina	234	225
ammoniaca?	ammoniaca metata	235	238
metata certe non aetites ?	not aetites	236	188
	=	237	160
filipes ? Iris	filopes Iris	238	161
amicta?	amicta, quite distinct from Iris	239	286
luteo-alba vitilis?	vitilis	240	189
? Omphalia tenuistipes? Omphalia mauretanica Maire?	tenella	241	190
acicula, mauvaise figure haematopoda	acicula haematopus (stem should be darker)	242	162
cruenta sanguinolenta, couleurs trop vives crocata	cruenta sanguinolenta	243	163
	galopus	244	207
galopoda epipterygia	epipterygia	245	208
? rorida	pelliculosa typical vulgaris	246	191
	plicato-crenata	247	248
rorida stylobates tenerrima	rorida stylobates tenerrima	248	249
Omphalia electica	discopus typical	249	192

No. of bound volume	No. printed on Plate		Соо	KE	Quiter
250	164	Agaricus	(Mycena)		
•		-	********	hiemalis	
251	193	a handere	Spiritone	setosus	
		********	Symmetric	capillaris	W/-
		#memory		juncicola	editor
252	239		(Omphalia)	hydrogrammus	non verdåtre
253	287	****	denidend	maurus	non verdany
		Section 1	page restri	offuciatus Postii	4.5 %
254	194.		Marie Co.	pyxidatus	
0.55	288	Market Market	-	leucophyllus	2
255	200	participal and the second	Open Services	striaepileus	
0.56	240	Annual	Name of the Control o	telmatiaeus	La St. Inc.
256	289	*******	West of	sphagnicola	
257	209		nentite	philonotis	leucophylla
258	209	bo trong	1 Species	oniscus	and the second s
230	209	, mentur	MAC V de	caespitosus	umbellsfera var. flava
259	250	Account	prom.d	demissus	No.
-39	1-00	to enterior	phy contri	hepaticus	Mr.
		***************************************	deces	muralis	pyxidata, male
260	271	refrance	PRODUCE	umbelliferus	
261	272	belon to	Water	buccinalis	21.5
	-/-	descriptions.	we-sering	retostus	cendré
		*******	range timper	abhorrens	- 344
262	241	Management	matter tout	pseudo-androsaceus	No. of
	1	Manageral	Merchan	griseo-pallidus	
	1 1 1	P *******	-	stellatus	Nissad
263	273	-	Septem Sept	campanella	w ev
-		-		- var. badipus	we code
		-	Mingration	pictus	*
264	210	******	N. artinus	camptophyllus	ers when
-		Process	Meth 14	griseus	v - 4×4
265	274	-	-	umbratilis	MAR OF
			A terr of cape	fibula	IF WF
266	251	********	nhor/Post	directus	*** ***
C		And-1940	WAVE A	belliae	rorida (vetustion)
267	252	*****	\$100-10 (m)	gracillimus	stellata on integrella
	1.	Bertina	pr = 40	bullula	M. Micheliana
~60	1 1	****	/ TV2	integrellus	brone is
268	290		(Pleurotus)		diyinus
269	226	-	process the eye	dryimis	10.00
270	253	-	No. ope	spongiosus	ostreatus var, columbinus
271	227	arbone.	Monate	ulmarius	8.01 1/2
272	254		Married .	tessulatus	AUG.
273 274	² 55 256	*******	Annual or an annua	subpalmatus	The second second
275	178		Baryaday	craspedius fimbriatus	ulmarius
-/3	1,0			Ruthae	palmatus
276	257			lignatilis	Normal District
. 70	737	Name of Street	-	circinatus	679439
277	179		*****	pantoleucus	conchatus
277 278	275		former	pantoleucus	CAMPENGENCY
	-,5		-	mutilus	scyphoides (major)
279	195	-	-	ostreatus	acypnomes (major)
279 280	195	-		- var. euosmus	Endow.
281	180	-	Married	revolutus	
282	228		-	salignus	ostreatus (genuinus)
283	291		-	acerinus	dryinus (genunus)
284	258	A	-	petaloides	weeken Sarakan
				serotinus	malè

Maire	REA	No. of bound volume	No. printed on Plate	
 corticola	corticola	250	164	
hiemalis?	hiemalis setosa	251	193	
capillaris	capillaris —			
maura	hydrogramma maura	252 253	239 287	
Postii	Postii pyxidata	254	194	
pyxidata ?		255	288	
		256 257	240 289	
onisca	onisca	258	209	
umbellifera var. flava — —	umbellifera var. flava — hepatica	259	250	
umbellifera	muralis umbellifera	260 261	271 272	
	retosta abhorrens —	262	241	
griseo-pallida ? campanella Marasmius fulvo-bulbillosus	stellata, but should be white campanella cauticinalis	263	273	
picta?	picta camptophylla	264	210	
grisea —	grisea umbratilis	265	274	
fibula et var. Swartzii	fibula et var. Swartzii pseudo-directa	266	251	
Mycena rorida? integrella?	 gracillima bullula	267	252	
cuspidata Quél.?		268	290	
corticatus = dryinus corticatus = dryinus	corticatus dryinus	269	226	
ostreatus var. columbinus	columbinus	270	253	
ulmarius	ulmarius	271	227	
wante		272	254	
palmatus	palmatus	273	255 256	
ulmarius	not craspedius	274	250	
lignatilis?	fimbriatus	275	178	
lignatilis	Ruthae lignatilis circinatus	276	257	
Panus torulosus?	palmatus spores wrong for pantoleucus	277 278	179 275	
mutilus (forme de Omphalia scyphoides) cornucopiae	mutilus sapidus (spores lilac in mass)	279	195	
ostreatus var.	sapidus	280	196	
Panus torulosus?	revolutus var. anglicus Massee	281	180	
ostreatus	ostreatus	282	228	
corticatus? anneau non visible	not acerinus	283	291	
geogenius forma petaloides serotinus	petaloides serotinus	284	258	

No. of bound volume	No. printed on Plate		Coo	KE	Quéter
285	211	Agaricus (Pleurotus)	mitis	May .
286 286	276	118000000	Aprend .	gadinoides	du tyorhizus
200	2/0	and the	#de-codes	limpidus	* X
			Manager 6	reniformis	12 mag
287	242	*******	Man to My	lauro-cerasi	*** F FM
,		8483	and the second	tremulus	male
		and 100 miles	0.000	acerasus	in alpha
288	259	-	w/10-175	porrigens	nakiri iy
		nutromité	previous	septicus	g ₁ ,····································
289	243	therees	accepted	mastrucatus	Montality
		Married .	8 0-140-140	atracoeruleus	algidus (junior)
290	260	-	per meth	Leightoni	male
			parent A	algidus	111411
291	244	******	substitute agranisms	fluxilis	TO 100
		*****		cyphellaeformis	pon! striatulus?
		Special St.	New yorks	applicatus Hobsoni	dictyorhizus
292	212		944F-109	striatulus	***************************************
	1 1	Managage .	\$400 MA	hypnophilus	1.454
				chioneus	**
000	000		(Valvaria)	bombyeinus	ARTES
293	293		(vorcania)	volvaceus	4 V=r
294	294	4000	arrest.	Loveianus	= fuberidus
295 296	295 296		Secretal Secretal Secretary	Taylori	1.8
297	297		****	speciosus	Carted*
298	298		simusia.	gloiocephalus	west
299	299	******	*******	medius	Acollogia
300	300		*******	temperatus	th radii
3	3		Name and Address of the Owner, where the Owner, which is the Owner, whi	parculus	In Found
301	301	_	(Pluteus)	cervinus	60
302	565	-	become	- var. patricius	APM-133
303	302		SUCCESSION .	- var. eximius	No. of
1				D1122	
304	357		Systema	- var. Bullii	Indiana
305	303		4404	- var. petasalus	hopfast
306	304	Section II	garring garring	umbrosus hispidulus	tous deux malé
307	517	*******	Parameter .	ephebius	malè
308	597		Printers	pellitus	pas la spore
309	305		N-MANUEL .	nanus	lutescens : chrysophaeus Q.
310	325	_	British -	spilopus	cervinus
311	518		governo.	semibulbasus	4.0np.w
				violarius	Russula lateritia on nitida
312	598		*******	roseo-albus	non
313	421		4 ·	leoninus	(Manage)
314	309		Yearing	chrysophaeus	cervinus (gracilis)
315	422	_	andrey	phlebophorus	rhodopolius, leoninus caa Pleur, palmatus
316	310		(Entoloma	simuatus	= lividus ou elpheatus major
	311	-	- Carrier and Assessment Con-	lividus	malè
317 318	469		*****	- var. roseus	potius clypeatus
319	312			prunuloides	and the state of t
320	313	E 1	-	repandus	Inocybe repanda (Bull.) Qué
321	314	-		placenta	elaphinum Fr.?
322	339	-	-	helodes	prunuloides (genuinus)
323	373	1		helodes var.	Inocybe caesariata

Maire	Rea	No. of bound volume	No. printed on Plate	
nitis	mitis	285 286	211	
lictyorhizus?	-	286	276	
	-			
ent-yell	· .	_		
*****	Process	287	242	
2				
icerosus	acerosus	-00	0.00	
porrigens	porrigens	288	259	
epticus	septicus	289	243	
nastrucatus atro-coeruleus	mastrucatus atro-coeruleus	209	443	
algidus	auro-coerateus	290	260	
algidus?	algidus	-30		
aiguas.	fluxilis	291	244	
cyphelliformis	cyphellaeformis			
striatulus	applicatus			
dictyorhizus		292	212	
striatulus	striatulus			
hypnophilus	hypnophilus			
chioneus	chioneus			
bombycina, non typique		293	293	
volvacea	volvacea	294	294	
Loveiana	Loveiana	295	295 296	
Taylori	Taylori	296	297	
speciosa	speciosa gloiocephala	297 298	298	
speciosa, forme foncée (= gloiocephala)	media	299	299	
	nteata	300	300	
parvula	parvula	"		
cervinus	cervinus, stem characters	301	301	
	poor			
cervinus var. patricius	cervinus var. patricius	302	565	
cervinus var. eximius	cervinus var .eximius, colour	303	302	
	exaggerated	201	0.57	
	Bullii, a distinct species	304	357 303	
rigens (Pers.) = salicinus Lange	cervinus var. petasatus	305	303	
umbrosus Fr. non Pers., mais arête noire des lamelles non figurée	umbrosus)	306	304	
semibulbosus? ou hispidulus?	hispidulus)]	"	
semioutoosus . Od nispidatus .		307	517	
		308	597	
nanus	nanus)	309	305	
nanus var. lutescens	lutescens	1		
?	spilopus	310	325	
semibulbosus?	**************************************	311	518	
Un Lepiota ou Entoloma?	-		508	
palmatus	palmatus	312	598	
leoninus	leoninus typical	313	421	
leoninus var. caloceps	, , , , , , , , , , , , , , , , , , ,	314	309	
? forme grêle de P. cervinus	phlebophorus)		1	
phlebophorus palmatus	palmatus	315	422	
clypeatum?	sinuatum	316	310	
lividum, mauvaise figure	lividum	317	311	
?		318	469	
prunuloides, peu typique	_	319	312	
	prunuloides poor	320	313	
		321	314	
prunuloides, forme foncée	4	322	339	

No. of bound volume	No. printed on Plate	,	Coo	KE	Quilet
324	315	Agaricus (E	ntoloma)	Persoonianus	Pluteus semihulbasus vas Lept sericella major?
905	326	armenia.	Suratrus	Batschianus	mudidus (gracilis)
325 326	327	******	wex 107	Bloxami	A 64.000A
327	328	mental	********	ardosiacus	or Meso
328	581	weeklik	\$1.145kg	liquescens	Psathyra sarcocephala avec spore de Rhodophyllus
329	341		APPROVAMENTS.	ameides	malé, il est livide
330	470	anning	satisticals	frumentaceus	Cortin, firmus?
331	306	anapetes .		Saundersii	dypeatum vac, albescens
332	316	-	periods.	fertilis	lividum, malė
333	317	www	-	jubatus	races
334	318		professed.	resutus	
		and	atter et ,	griseocyaneus	male
335	307		promise of	sericellus	treates .
335 336	374		protects	Thomsoni	scabiosum ou griseo-cyaneum
337	319	****	******	clypeus	clypeatum Fr.
337 338	342		dimension .	rhodopolius	clypeatum
339	329	-	-	Wynnei	Pl. villosus
340	320		Provide Co.	costatus	Howard
×		-	Principal	sericeus	Pin ne
341	321	******	Property	nidorosus	survey.
342	308		water	speculum	Plut. pellitus, non la spore
343	322	- (0	Ilitopilus) prunulus	trop blanc
344	323	******	Million W	orcella	Membe
345	375		Non-resp	mundulus	pas blanc
			directed.	cretatus	O. albula?
346	485		Minerary	popinalis	_desired
347	486		Management of the last	undatus	No. of a contract of the contr
348	501		Ministrati	cancrinus	reproduce
349	324	,	*******	carneo-albus	No. Asser
		Massa	******	stilbocephalus	Nolanea incarnata
350	599		Minester 8	stilbocephalus var.	Coll. laccata
351	487		Military	vilis	Lept. chalybaea (vetusta)
352	330	(.	Leptonia)	placidus	Mr Hore
353	331		direction of	lampropus	aling-task
354	332		en-	aethiops	rimany
			****	solstitialis	NOCONN
355	333		enne.	serrulatus	anatina?
356	334		Services .	euchrous	simula
357	335		******	chalybeus	4108,36
357 358	549	***************************************	Photos	lazulinus	No rough
359	336		Amour	incanus	euchlorus
360	488		Printer	formosus var, suavis	? malé
361	337		***	chloropolius	Nolanea icterina?
362	376	- (.	Nolanea)	pascuus	Moneya
363	377	-	-	Babingtonii	pas blanc
· 4			*******	mammosus	ferror
364	378	_	The	pisciodorus	Analoge
× 2 -			~	rufo-carneus	Lept. solstitialis ou C. laccate
365	338		- Think	icterinus	Name
366	379	_	******	piceus	Yelephon
	1		Arrens	coelestinus	?
367	340		-	verecundus	Weeke
	-	-	-	rubidus	Leptonia sericella
	1 0	1.			
368	380	- /	Eccilia)	parkensis	

Maire	Rea	No. of bound volume	No. printed on Plate
MAN	Providence (324	315
gran-mild		325	326
Bloxamii = madidum	Bloxamii	326	327
Eccilia Mougeotii forma leptonidea	ardosiacum typical	327	328
L'hypothèse de Quélet est très vrai- semblable		328	581
ameides	ameides	329	341
Inocybe jurana Pat.=I. frumentacea Bres. (an Bull?)	rhodiola	330	470
Lini Jam	Saundersii	331	306
lividum	porphyrophaeum	332	316
porphyrophaeum	porproyropaucum	333 334	317 318
griseocyaneum, mauvaise figure	griseocyaneum	334	3.0
Lept. sericella	sericella	335	307
? ^	-	336	374
clypeatum Fr.	clypeatum, rather poor	337	319
rhodopolium Fr.?	rhodopolium	338	342
Energies.		339	329
costatum	costatum sericeum	340	320
sericeum nidorosum	nidorosum	341	321
ntaorosum speculum	speculum	342	308
prunulus	prunulus	343	322
prunulus âgé	prunulus	344	323
pas mundulus	not mundulus	345	375
cretatus	cretatus		*
	popinalis	346	485
? teinte de Collybia nitellina		347 348	486
Eccilia cancrina	cancrinus sericella		501
Leptonia sericella?	stilbocephala	349	324
Nolanea incarnata? Laccaria laccata?	Smithii Massee	350	599
Eccilia Mougeotii?	vilis	351	487
placida	placida	352	330
lampropoda	lampropus	353	331
_ * *	aethiops	354	332
	solstitialis		
serrulata var. Berkeleyi Maire	serrulata var. Berkeleyi	355	333
euchroa	euchroa, larger than usual	356	334
chalybea	chalybea lazulina poor	357 358	335
incana = euchlora	incana	359	549 336
		360	488
Nolanea icterina?		361	337
staurospora Bres. à spores mal figurées	proletaria	362	376
Babingtonii (trop pâle)	-	363	377
mammosa	mammosa	-6	0
Naucoria Cucumis	Cucumis	364	378
Nolanea infula?	rufo-carnea	26=	008
icterina	icterina Cucumis	365 366	338
Naucoria Cucumis	Gucunus	300	379
<u>.</u>	verecunda	367	340
Clitopilus cretatus? spores non angu-	rubida	3-1	3.1
leuses	*		÷ 1,
		368	380
*****	carneo-grisea	1	1

No. of bound volume	No. printed on Plate	Со	OKE	Quélet
369	613	Agaricus (Eccilia)		malė
	1	service discretiv	flosculus	
		on a cody.	acus	sericellus
370	343	Medical Medical	atro-punctus	Total Control
		101 1	rhodocyclus	Welling
371	344	— (Claudopus)		
		Administration of the second o	depluens	pas blanc
			byssisedus	blane
372	345	- (Acetabularia		Hyphol, fatuara, Psathyra torjens
373	346	(Pholiota)	aureus	Ministra
374	347	45 mining lander lander	- var. Here-	benč, genomus
			fordensis	
375	348	annes, proper	caperatus	malé, plutôt aureus
376	349	Virgon exists	terrigenus	So was
377	358	Mretina phonicis	erebius	pas violeté
377 378	359	Market made control of	ombrophilus	errhia?
379	350	Mingleson Michigan Market Company	togularis	(mm Bull.) Fr. = Arrhenii F
380	423	#6 rywyd darfa writh	durus	très blanc!
38r	360	Meditology (Mondaying)	praecox	2
. •	1			•
382	361	Mineral errorsh	radicosus	the diagraph
383	362	Mining market	pudicus	61 M.1 W
384	363	Arresty process	leochromus	aegerita (gracilis)
385 386	364	Maryland Committee Committ	capistratus	aegerila (vetustior)
386	453	before (4)	aegerita	male, trop jaune
387 388	365	mention weather	aegerita	Ale Charte
388	600	Triprotes desiring	comosus	= destruens Brond.
389	366	Whethe	heteroclitus	destruens
390	351	Printers - Statement	aurivellus	
391	367	Manager Malacra	squarrosus	tenè
392	614	personal mensonal	- var. Mulleri	# **vident
393	614	- many	- var. verrucu-	Mortopia -
	1 1		losus	
394	352	harries same	spectabilis	aureus
395	353	President	adiposus	1964/1/14
396	368	- American	flammans	malè
397	369	distribution personal	junonius	?-aurea (gracilior)
398	370	Afficiation.	tuberculosus	**************************************
200		Minimum Minimum Minimum	curvipes	Arrage 2
399	502	Mining Mining	cruentatus	ons Flammula azyma Fr.?
400	371	Profession colonials	dissimulans	Ph. erebia
401	354	nation (All)	Cookei	ou gummosa
402	355	Meaning - Meaning	mutabilis	Walding
403	372	Monthly	marginatus	tanicalor
404	356	Marine Marine	mustelinus	4(Linguage
405	500		unicolor	marginata
4-0	503	Marine William	pumilus	var. humicola
406	424	17	mycenoïdes	?
407	389	- (Inocybe)	hystrix	Attorney
408		Protes	calamistratus	With respect
100	582	- Annual	lanuginosus	dulcamara
409	425		dulcamarus	lucifuga
1-3	4-5	(Inocybe)	plumosus	**************************************
410	390	- Marine	cincinnatus	Minny
	472		haemactus	corydalina (vetustior)
411			pyriodorus	
411 412	473		incarnatus	l caesariata repanda

Maire	Rea	No. of bound volume	No. printed on Plate	
?	griseo-rubella flosculus	369	613	
Leptonia sericella (forme décurrente)	acus atro-puncta	370	343	
variabilis	variabilis	371	344	
byssisedus (trop gris) Bolbitius bulbillosus	depluens byssisedus acetabulosa	372	345	
spectabilis	aurea	373	346	
aurea Fr. non Quél., forme colorée	aurea	374	347	
aurea Fr. non Quél. Flammula gummosa	ochrochlora typical	375 376	348 349	
erebia (trop violacé)	erebia		358	
Peut-être un gros Stropharia inuncta?	ombrophila, very large ex- ample	377 378	359	
togularis Fr. non Ricken	togularis dura	379 380	350	
dura (trop coloré) praecox (fig. sup.)		- 2	423	
dura (fig. inf.)	praecox	381	360	
Hebeloma radicosum	radicosum	382	361	
cylindracea = aegerita	aegerita	383	362	
cylindracea	leochroma	384	363	
cylindracea forma fuscescens	aegerita	385 386	364	
cylindracea forma fuscescens	aegerita aegerita	387	453 365	
cylindracea forma destruens	destruens	388	600	
destruens forma	destruens	389	366	
aurivella ou adiposa?	aurivella	390	351	
squarrosa	squarrosa	391	367	
Mulleri (espèce distincte)	Mulleri	392		
squarrosa (forme pâle)	squarrosa var. verruculosa	393	614	
spectabilis	spectabilis	394	352	
adiposa	adiposa	395	353	
flammans	flammans	396	368	
spectabilis, forme grêle	Junonia	397	369	
t curvipes	squarrosa var. verruculosa curvipes (flesh of stem darker)	398	370 502	
	cruentata dissimulans	399 400	371	
ombrophila?	ochrochlora	401	354	
Flammula gummosa mutabilis	mutabilis	402	355	
unicolor	unicolor	403	372	
CV PANA PACE	mustelina	404	356	
marginata	marginata	77.0		
marginata	pumila	405	503	
?	togularis			
hystrix	hystrix	4.06	424	
calamistrata	calamistrata	407	389	
dulcamara?	dulcamara	408	582	
lucifuga ?	dulcamara plumosa prob., but no spores	409	425	
lanuginosa	lanuginosa haemacta	410	390	
piriodora ssp. haemacta	pyriodora	411	472	
piriodora	incarnata	412	473	
jurana Pat.	scabra	413	391	

No. of bound volume	No. printed on Plate		Coc	OKE	Quélet
414	392	Agaricus	(Inocybe)	maritimus	lanuginosa
415	583		~~~	lacerus	Manager Annual Control of the Contro
416	393	-	Server Miles	flocculosus	rimosus (gracilis)
417	381	***************************************	Monthly	Bongardii	rimosu
417 418	382	an-orași.	encentery.	muticus	. 1109
419	426	*****	advant-us	carptus	forma robusta: hirsuta?
420	394	personal	Merchany	deglubens	caesariata
- "	1			obscuries	Johnana
421	427		-		cincinnata
422	395	-	Residence	echinatus	Inacybe??
423	504	******	Annual State of State	schistus	rimosa on fastigiata
424	454	***************************************	-	fibrosus	benë
425 426	396	special	- House	phaeocephalus	rimosa vel brunnea
426	383	a decidade	WHITEHOUSE	fastigiatus	tomentosa
427	397	******	Menty	hiulcus	repanda Bull.
428	398	*******	Policy	Curreyi	caesariata?
429	384	Medicality	#*************************************	rimosus	workaye
430	385	******	Willeria	asterosporus	Ziroliba
431	386	-	\$1000m	eutheles	being the
432	505		Warrang	margaritispora	? lacera (vetustissima) pas l spore
433	387	*********	/*******	destrictus	Moderation
434	519	********	Wrong	perbrevis	Cortinarius incisus Fr.
435	428		**********	descissus	returns .
			Prin sang	Trinii	scabella
436	399	-		sambucinus	probany
437 438	388		erroring protection	caesariatus	capucina Fr.
430	400	*******	distributes.	sindonius	geophila (luxurians)
439	429	-	Miningray	lucifugus	male
110	401		Trebute	Clarkii	geophila on tomentosa
440	402		Windows.	geophyllus scabellus	fraction ()
441 442	520		-	Rennyi	male
443	403	Second Second Second Problem	Property	trechysporus	fusca?
TTO	4-2		Marine	vatricosus	plumosus?
444	404		-	Whitei	tomentosus
TTT	ToT	******	-	tricholoma	lucifuga?
445	405	(Hebeloma)		- which
445 446	406	· · · · · ·	- Acoctonica)	fastibilis	Mydelex
11.	4			Justones	- Hondon A
447	407	Ministra	Persons	senescens	
448	430	Reference.	Melvining	glutinosus	
449	408	-	Military	testaceus	>
				***************************************	(forme de crustuliniformis
450	409	******		firmus	aspect de Cortinarius
				J*************************************	imbutus on varius
451	410	-	-	claviceps	persibellis
452	411	_	Wilnes	mesophaeus	versipellis
453	412		PPANNS	- forma minor	2
454	506		-	subcollariatus	· ·
455	413	******	M-MINA	sinapizans	versipellis
455 456	507	-	-	crustuliniformis	crustuliniformis B.
457	414	-	-	- forma minor	Minima
457 458	415		Manage	longicaudus	elatus
459 460	416		mentage menta	- var. radicatus	versipellis
460	417	-	-	truncatus	wer aspected a
10 10 10 10 10					
461	418	-	-	nudibes	elatus
462	419			capniocephalus	versipellis

Maire	Rea	No. of bound volume	No. printed on Plate	
? ? non lacera Ricken	maritima —	414 415	392 583	
?	flocculosa	416	393	
? non Bongardii	not Bongardii	417	381	
2	mutica	418 419	382 426	
P	deglubens	420	394	
obscura	-)		1	
cincinnata?	<u> </u>	421	427	
Lepiota echinata	Lepiota haematosperma	422	395	
? ? Entoloma sp.?	schista fibrosa	423	504	
maculata Boud.?	phaeocephala	424 425	454 396	
fastigiata		426	383	
?	Godeyi	427	397	
?	fastigiata	428	398	
Queletii?	probably Queletii	429	384	
asterospora = rimosa Fr. genuina eutheles	asterospora tomentosa	430 431	385 386	
eunetes .	margaritispora	432	505	
	destricta	433	387	
non perbrevis. Cort. incisus?	_	434	519	
r anghalla	Trinii	435	428	
scabella	sambucina	436	300	
Access 1	caesariata		399 388	
geophylla var. alba?		437 438	400	
lucifuga	lucifuga	439	429	
geophylla var. alba? geophylla var. alba et violacea	geophylla	440	401	
3.7.5	-	441	402	
3		442	520	
trechispora	trechyspora	443	403	
		444	404	
Ripartites Tricholoma	tricholoma	444	404	
		445	405	
fastibile?	fastibile, pileus too deep in colour	446	406	
senescens	senescens	447	407	
Flammula lenta	glutinosum, pileus too dark	448	430 408	
? pied ordinairement bulbeux	testaceum	449	400	
5	_	450	409	
versipelle var.		451	410	
fig. sup. versipelle	versipelle	452	411	
fig. inf. versipelle var. mesophaeum	mesophaeum \(mesophaeum \)	453	412	
versipelle var. mesophaeum		454	506	
sinapizans	sinapizans	455	413	
sinapizans forma	crustuliniforme	456	507	
crustuliniforme	crustuliniforme var. minus	457	414	
longicaudum?	longicaudum	458	415	
Phodobarillus trameatus ? ? enores tron	radicatum Tr. truncatum	459 460	416	
Rhodopaxillus truncatus??spores trop grandes et trop amygdaliformes	11.41414444116			
	(; - · · · · · · · · · · · · · · · · · ·	461	418	
versipelle var. mesophaeum	mesophaeum	462	419	

No. of bound volume	No. printed on Plate	Соон	CE	Quérer
463	420	Agaricus (Hebeloma)	ischnostylus	elatus (gracilis) ou saccha-
464	508		magnimamnu	serripellis
465	431	— (Flammula)	petiginasus gymnopodius	non't mesophaeum I male
466	437	Transpara students	vinosus	pas la couleur, Plear, Espagn?
467	438		floccifer	Heb. glutmosum
468 469	500 439	Managan Angarigas	decipiens clitopilus lentus — long-stemmed	Tricholoma truncation?
470	440		form	*
471 472 473 474 475 476 477 478	474 475 441 476 442 432 433 434	Parling Microso, Simple Application Simple State of the S	mixtus juncinus gummosus spumosus carbonarius fitius	carbonarius almicola spumaca? ? ? Gartinarius Heli, senescens Pholiota aurea
479 480 481 482 483 484 485 486 487 488 489	435 443 444 477 445 446 436 615 447 448 616 449	British and a service of the service	astragalinus alnicola flavidus inauratus conissans inopus apicreus hybridus sapineus picreus ochrochlorus helomorphus	mulė spunosus hybrida ?? malė Pholiota confragosu liquiritiae gumnosa
491 492 493 494 495	450 451 452 455 601	- (Naucoria)	seambus filiceus cidaris Cucumis anguineus centunculus horizontalis	? graminis male copiés de Fries
496	509	process against	semiflexus rimulineola	horizontalis
497 498	456 489	and the same of th	rubricatus abstrusus innocuus	Marasmius ramealis pellucida Flam, carbonaria
499	457	0 8 1	cerodes melinoides	Pluteolus dictyotus Kalch.
500	490	printing guiden	pusiolus nuceus	malè
501	491		glandiformis badipes	containing and a second of the
502 503	478 458	Africana despitores Administra	scolecimus striaepes sideroides	Pluteolus dictyotus
504	617		triscopus vervacti	male
505	492		tenax pediades	autochthona

Maire	Rea	No. of bound volume	No. printed on Plate
sacchariolens?	sacchariolens	463	420
Inocybe sp. Inocybe fastigiata? forma		464	508
Clitocybe olearia? mais chair trop blanche	sapinea	465	431
Pleurotus Eryngii?		466	437
lenta ?	decipiens	467	438
? lenta lenta	glutinosum glutinosum	468 469 470	500 439 440
carbonaria (grêle) alnicola?	carbonaria	471 472 473	474 475 441
carbonaria ? H. senescens?	carbonaria, large form	474 475 476 477	476 442 432 433
forme décurrente stérile de <i>Pholiota</i> spectabilis ? rubicundula Rea	rubicundula	478	434 435
alnicola	alnicola flavida inaurata	479 480 481 482	443 444 477
conissans, mauvaise figure Hypholoma epixanthum Fr.? Quél.! — Ph. confragosa	conissans Hypholoma radicosum Lange apicrea confragosa	483 484 485 486	445 446 436 615
sapinea ?	sapinea ochrochlora	487 488 489	447 448 616
? Ripartites Tricholoma	helomorphus, spore wrong scamba	490	449
Cucumis d'après Fries centunculus, trop pâle	Cucumis anguinea centunculus	491 492 493 494 495	450 451 452 455 601
horizontalis —- rimulincola	horizontalis	496	509
Mar. ramealis ? Flam. carbonaria	ramealis furfuracea	497 498	456 489
	cerodes melinoides pusiola	499	457
? Cortinarius		500	490
badipes	badipes escharoides	501	491
striipes	striaepes —	502 503	478 458
Galera triscopa vervacti	vervacti	504	617
? n'a pas la teinte de N. autochthona pediades	pediades	505	492

No. of bound volume	No. printed on Plate		Coo	KE	Quilet
506	479	Agaricus	(Naucoria)	arvalis	and S pain
		115011000	(21200110)	semiorbicularis	- Manager
507	493	on-rentered	William Ave.	tabacinus	
508	494	-	estion of	myosotis var. major .	nade, Cortinarius
509	459	annote the same of	mirror.	temulentus	male
3 3	1				
510	482	***	PP-998	latissimus	Goll, extuberans? Ps. spadices
511	510	merchan	dividional	porriginosus	Cartin, scandens
512	511	*******	en-rises.	sobrius	crobulus
		*******	Freedy	- var, dispersus	# scople
513	480	*****	provide	erinaceus	The Property of the Control of the C
		********	*****	siparius	? il est humicole
514	512	day	lates months	conspersus	malė
		White	20-20-4	escharoides	malé
515	513	******	Th-wife.	carpophilus	Delines
			Fall I N	graminicola	inquilina
516	495	(Pluteolus)	reticulatus	PR PR CALL
517 518	460	•	(Galera)	lateritius	Bolbitius apalus
518	461	-	discount .	tener	re Ina
		amban.	perception	- var. pilosellus	11 1996
519	462	-	(Princip)	ovalis	tenera
520	463	timetuna	Processo	antifus	tenera
	.0.	-	arcenda provend Monada	confertus	- Maria Sapal
521	481	annual of	Mrinorada	sparteus	Michaele Programme Control of the Co
	.6.	-	This work	pygmaeo-affinis	
522	464	Minima	Military .	vittaeformis	male
F00	46-	Adhiyas	MOVE AND ADDRESS OF THE PARTY O	rubiginosus	deposits. delegé tos
523	465	*******		hypnorum	
504	466	-	and the same of th	hypnorum	1 %
524	400		territorius.	mniophilus minutus	malė
525	467		The second	ravidus	est humicole malé
3-3	407			mycenopsis	
526	602	(Tubaria)	cupularis (var.)	Myc. epipterygia pellurida
	603			furfuraceus	persuctura
527 528	483		*****	- var. trigono-	
5	1-3			phyllus	V-1704
529	484		Allerening	paludosus	Alphoto-yag
530	468	******	THINNS	stagninus	emaging
531	514	-	amening.	embolus	Alle or sea
00		*****	Rivolog	autochthonus	undulatus Bull.
532	496	******	767-9786	crobulus	***************************************
533	497	*******	Process	inquilinus	malè
534	499	((Grepidotus)	alveolus	Martina
		`	-	calolepis	J15ppin
535	498	-	arresa	mollis	W1500W8
535 536	515		******	haustellaris	0.719.86
		-	ammonia Malicon Moure (a)	Rubi	relam on
		-	MATRICE .	Phillipsii	Affilia da Affil
	1 1	princes	Mirana	chimonophilus	variabilis
537	516		******	epigaeus	applanatus
		- minut	Printer	Ralfsii	esser.
		ķ	Seems .	epibryus	Milyste
0			-	pezizoides	Pleurot, craterellus
538	521	(Psalliota)	augustus	villatica ?
539	522			Elvensis	Alaba.
540	523		-	arvensis	eMinus.
541	584	1 -	-	- var. purpur-	sylvatica var. rubella Gill.
542	524	1		ascens	
J44	344	-	*******	cretaceus	Lepiota pudica

Maire	Rea	No. of bound volume	No. printed on Plate
arvalis	arvalis	506	479
semiorbicularis tabacina ?	semiorbicularis tabacina	507	493
hrmaqua.	Myosotis typical	508	494
Tubaria pellucida? mais spores trop grandes, et voile non figuré	temulenta	509	459
?	Miniminal	510	482
i a de la companya de		511	510
Tubaria crobulus	sobria pusiola	512	511
erinacea	erinacea	513	480
potius Lepiota conspersa	siparia conspersa	574	512
escharoides?? spore trop allongée	not escharoides	514	312
carpophila	carpophila	515	513
Tubaria inquilina?	grāminicola		
reticulatus	reticulatus	516	495
lateritia		517	460
tenera	tenera	518	461
e grege G. tenerae tenera	pilosella ovalis	519	462
antipoda	antipus	520	463
spartea pygmaeo-affinis	spartea pygmaeo-affinis	521	481
vittiformis? teintes du chapeau fausses		522	464
_	rubiginosa	3	1 1
Hypnorum Sphagnorum	hypnorum	523	465
lateritia?		524	466
inquilina?		505	467
Myc. epipterygia?	not mycenopsis	525	407
pellucida	pellucida	526	602
furfuracea	furfuracea	527	603
furfuracea forma	furfuracea var .trigonophylla	528	483
paludosa	paludosa	529	484
stagnina	stagnina	530	468
		53 I	514
autochthona	autochthona		6
crobulus	crobulus inquilina	532	496
inquilina ? palmatus ?	alveolus	533 534	497 499
calolepis	calolepis	334	433
mollis	mollis	535 536	498 515
Nauc. effugiens Phillipsii	effugiens Phillipsii	332	
applanatus Quél. (an Fr.?)	Dalfaii	537	516
	Ralfsii		
n'est pas P. craterellus			
Agaricus augustus	Psaliota augusta	538	521
Elvensis	Elvensis	539	522
arvensis Fr., Quél., non Pat. silvicola Vitt. var. purpurascens Cke.	arvensis sylvicola var. purpurascens	540 541	523 584
xanthodermus Genev. var. lepiotoides Maire ou Lep. naucina	lepiotoides Maire	542	524

No. of bound volume	No. printed on Plate	Co	OOKE -	Quint
543	FOE	Agaricus (Psalliota	i) bratensis	
544	525 526	218011000 (1 3001000	campestris	sillatica Brond.
		Brown by the self-to-	var dontensis	ellatica
545	527	Brown by St. 14-36		
546	528	Sections section	- var. costatus	champignons alteres
		British Physics	var. exannu- latus	villation
547	529	Profession was a second	- var. sylvicola	sylvatica, male
547 548	585	Shropped Hirotopy	- var. villaticus	Pilot 144 144 14 1 X 1 A 1 A 1 A 1 A 1 A 1 A 1 A 1 A 1 A
549	530	Many - serveral	sylvaticus	malé
JTJ	33.	Propose American	sylvaticus	111441
550	531	months of the contract of the	haemorrhoidarius	
551		- Phones Million	subgibbosus (var.)	Distinct
	532		comtulus	Pholiota negerita, vetustio
552	533 618	(Dilagge)		?
553	010	- (Pilosace)	Algeriensis	•
554	550	- (Stropharia)	Percevali	Flammula spumosa
555	551	Managatia Philosoph	aeruginosus	
555 556	552	gimbolog Managag	albo-cyaneus	
557		inneres enteres		aeruginosa
	534	months and and	inunctus	male
558	535		coronillus	lamelles brun-paurpre
559 560	536	Military Museumpe	melaspermus	benic
500	553	Herman Various	squamosus	
561	554	provided with date of	thraustus	MAT A &
562	555	distribuyer of hamilton	- var. auranti-	? etuetorum
563	556	Market Market	Worthingtoni	deruginosa
564	604	teriporas (constant)	luteo-nitens	Hyph, appendiculatum Bul
565	537	Marine Marine	merdarius	Flammula carbonaria
566	538	etwee strong	stercorarius	semiglobata
567 568	539	Minute Amount	semi-globatus	SETTING COLUMN
568	540	manage manages	caput-medusae	? colonea, vetustion
569	541	Annage Andreas	Jerdoni	
570		describe project	spintriger	здиатога
571	542 619	And house Substitute	hypsipus	11 . 1
572	557	- (Hypholoma)	nypsipus enhlatorition	Hyph. Candolleanum
573	558	(11) protonta	sublateritius	faccine
574	559	endrous Phones	- var. squamosus	- Harris
575	560	Province Province	capnoides	Market
575 576	561	Production Branching	epixanthus	4097014
577	562	destroy despess	fascicularis	Anna San San San San San San San San San
578	586	Property Services	- var. elaendes	Andrew to
579	587	Property Briens	dispersus	malė
3/9	507		oedipus	Stroph, versicolor
580	740	Markey Kindrey	punctulatus	was sales
	543	Median William	storea var. caespi- tosus	Stroph, cotonea
581	566	Personal marriages	lacrymabundus	? cotonea, vetustior
582	563	Personal Continues	velutinus	
583	564	Annable Services	pyrotrichus	Indoor
584	544	-	cascus	appendiculotum
585 586	545 546		lanaribes	
500	546	-	Candolleanus	appendiculatum, vetustius
587	547	-	appendiculatus	appendiculatum malè
-				
588	548 605	-	leucotephrus	
589	605	-	egenulus	appendiculatum
100				Psathyra cermua
590	567	- (Psilocybe)	hydrophilus	malè

Maire	Rea	No. of bound volume	No. printed on Plate	
pratensis?	pratensis	543	525	
campestris	campestris	544	526	
villaticus?	hortensis as species	545	527	
haemorrhoidarius? forme pâle				
forme de pratensis?	varieties of campestris	546	528	
silvicola? pas assez blanc	sylvicola as species	547	529	
villaticus	villatica as species	548	585	
silvaticus?	sylvatica	549	530	
silvicola var.	perrara			
haemorrhoidarius	haemorrhoidaria	550	531	
Ph. cylindracea (= aegerita)	. 7	55 I	532	
comtulus	comtula	552	533 618	
forme à très grandes spores qui reste	holisons	553	618	
forme de Stroph, depilata Fr. (teste	Percevalii, distinct from	554	550	
Plowright)	depilata			
aeruginosa	aeruginosa	555	551	
aeruginosa var. albo-cyanea	albocyanea	556	552	
inuncta	inuncta	557	534	
coronilla	melasperma var. lutescens	558	535	
melasperma	melasperma	559	536	
squamosa	squamosa	560	553	
squamosa ayant perdu les squamules du chapeau?	squamosa var. thrausta	561	554	
?	squamosa var. aurantiaca	562	555	
aeruginosa var. Worthingtonii	albo-cyanea	563 564	556 604	
merdaria?	merdaria	565	537	
stercoraria = semiglobata	stercoraria	566	538	
stercoraria = semiglobata	semiglobata	567	539	
caput-medusae	caput-Medusae	568	540	
Hyph, lacrimabundum Fr.?	-	569	541	
spintrigera	spintrigera	570	542	
?	hypsipus	571	619	
sublateritium	sublateritium	572	557	
sublateritium forma squamosum	sublateritium var squamosum	573	558	
capnoides	capnoides	574	559	
fasciculare?	epixanthum	575	560	
fasciculare	fasciculare	576	561	
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	not elaeodes	577 578	562	
dispersum peu typique melantinum?	not dispersum		586	
metantinum:	_	579	587	
lacrimabundum Fr. non Quél. nec Bull.	lacrymabundum Fr.	580	543	
idem	caput, Medusae	581	566	
velutinum	velutinum	582	563	
velutinum var. pyrotrichum	pyrotrichum	583	564	
Candolleanum	cascum	584	544	
Candolleanum forma?	appendiculatum	585	545	
Candolleanum	at the disculations	586	546	
Candolleanum, mais la forme grêle cespiteuse à gauche est peut-être	appendiculatum	587	547	
P. cernua leucotephrum, me paraît distinct	leucotephrum	588	F 48	
Candolleanum?	ocaurectus aus	589	548 605	
		209	005	
hydrophilum, mauvaise figure	-		100	

No. of bound volume	No. printed on Plate		Coc	KE	Quélet
591	620	Agaricus	(Psilocybe)	sarcocephalus	*-petition/
592	568			ericaeûs	
593	588	Acres 100	Proceed	sub-ericaeus	
594	569	Newscools		udus	
595	621		and the same	canofaciens	Rongel
596	570			areolatus	Fl. epixantha
597	570 622	-	-	agrarius	Mycena sudora
598	607		******	scobicula	Hyph. appendiculatum
599	606		-	chondrodermus	SAMOTOR
-				ammophilus	Cort. milvinus
600	608		-	coprophilus	malè
		Collector		bullaceus	malè
60 I	609	-	-	physaloïdes	malè, pas carminé
	=			nucisedus	Tub. inquilina
602	571	-	*****	atrorufus	male; Psathyra corrugis?
603	571 589	-	personal.	comptus	Pluteolus apalus
. •				hebes	nrane.
604	572		-	semilanceatus	New Property Company of the Company
60Ŝ	573	*****		- var. coeru-	
				lescens	
606	610		-	spadiceus	
607	574			cernuus	
6o8	590	-		foenisecii	
6		,	D	6.27	D
609	575	(Psathyra)	conopileus	Panaeolus fimiputris
610	591	***************************************	******	mastiger	Cort. acutus
611		P) Property		glareosus	Mycena zephirus
011	576			corrugis	Hyph. appendiculatum
612	592		***************************************	- var. vinosus	Hyph. appendiculatum
613	577	-	*****	pellospermus	Nolanea incarnata
614	577 611	Western	-	spadiceo-griseus	Hypholoma
	593		-	obtusatus	Hyph. fibrillosum
615	594		-	bifrons	Pan. acuminatus
777	331			var. semitinctus	Psathyrella gracilis
617	578	-	arreste .	semivestitus	
617 618	595			fatuus	malè
	000			fibrillosus	
619	579	******		helobius	fatua
620	579 580	******	Proper	Gordoni	pennata
		Manage	-	pennatus	
621	612	Manager Communication of the C		gossypinus	
		-		noli-tangere	-
622	596			microrhizus	bifrons
11	1.0		-	urticaecola	gyroflexa, minor
623	623	- (Pe	anaeolus)	separatus	malè
624	624		manage of the last	egregius	Strop. lacrymabunda Bull.
625	927		*****	leucophanes	separatus
		-		scitulus	Copr. ephemeroides
626	625			fimiputris	campanulatus
627	626	-		phalenarum	
628	627	******		retirugis	_
629	628		-	sphinctrinus	
630	629			campanulatus	conocephalus
631	630		_	papilionaceus	THE MANAGEMENT AND ADDRESS OF THE PARTY AND AD
632	631			caliginosus	acuminatus
			-	subbalteatus	
633	632	-	-	acuminatus	malè
		- X		fimicola	acuminatus
634	633	/ Da	.7 77 4	subatratus	

MAIRE	Rea	No. of bound volume	No. printed on Plate	
sarcocephala var. spadicea			600	
ericaea	ericaea	591	620	
sub-ericaea?	subericaea	592	568	
Hyph. elongatum (Fr.) Ricken?	elongata	593	588	
Psathyra helobia?	cionguia	594	569 621	
?		595		
?		596	570 622	
Hyph. Candolleanum forma?		597		
?		598	607 606	
ammophila	ammophila	599	000	
coprophila	coprophila	600	608	
bullacea	bullacea	000	000	
physaloides?	physaloides	601	609	
inquilina?	inquilina	001	009	
corrugis ?	_	602	571	
Galera apala?	-	603	571 589	
The star	hebes	003	209	
semilanceata	semilanceata	604	572	
ft wassey	var. coerulescens	605	573	
		505	5/3	
H. hydrophilum	hydrophilum	606	610	
cernua?	cernua	607	574	
foenisecii	foenisecii, typical except	608	590	
	spores special except	000	390	
Psathyrella subatrata?	conopilea	609	575	
Cort. acutus?		610	575	
Myc. zephirus?	Name of the last o	010	591	
P. conopilea? spores trop grandes	corrugis	611	576	
pour H. appendiculatum	corragis	011	5/0	
Hyph. Candolleanum		612	E00	
P. fatua?	Manusian.	613	592	
2	Williams	614	577 611	
3	Manage	615	593	
?	bifrons	616		
gracilis	var. semitincta	0.0	594	
semivestita	semivestita	617	578	
?	fatua	618	595	
?	fibrillosa	0.0	293	
helobia? pas typique	J	619	570	
S trans of transfers	Principal	620	579 580	
bennata	pennata	020	550	
gossypina?	gossypina	621	612	
P NES	Sandlana	021		
Psathyrella caudata?	-	622	596	
P. gyroflexa?	urticaecola	022	290	
Anellaria separata	separata	623	623	
Hyph. velutinum Fr.		624	624	
Anellaria separata?	*	625	927	
Copr. ephemeroides?		025	947	
campanulatus	fimiputris	626	625	
fimiputris?	campanulatus	627	626	
retirugis	retirugis	628	627	
phinetrinus	sphinctrinus	629	628	
campanulatus	campanulatus	630	629	
habilianareus	papilionaceus	631	630	
spores plutôt de campanulatus	campanulatus, small form	632	631	
2	campanana, sinan ioili	032	~y1	
· ·		620	632	
acuminatus	not fimicola	633	032	
subatrata	subatrata	634	633	
mentari 1666	1 Savariana	1 034	033	

No. of bound volume	No. printed on Plate	Сооке	Quélet
635	634	Agaricus (Psathyrella) gracilis	Psathyra corrugis
635 636	635	— hiascens	malè
637	636	aratus	Psathyra conopilea
637 638	655	trepidus	9
-3-	-55	- hydrophorus	
639	637	— caudatus	description
640	656	- pronus	crenata
•		pronus	- Angel ser
641	657	— empyreumaticus	Nauc. temulenta
• **	0.	— disseminatus	PUMPER
642	638	— atomatus	corrugis
-			gracilis
643	847	- crenatus	impatiens?
644	847 658	Coprinus comatus	Signal, SMM
645	659	ovatus	growth and the second
646	660	- sterquilinus	malė
647 648	661	oblectus	?
648	662	— atramentarius	Name - A four
649	848	- soboliferus	forme de fuscescens
650	663	— fuscescens	hydrophorus on micaceus
651	664	— var. rimoso-squamosus	deliquescens Bull.
652	665	picaceus	detamone
653	666	- aphthosus	Hyph, appendiculatum
654	667	— flocculosus	H. Candolleanum
655	668	extinctorius	***
655 656	669	— fimetarius var. pullatus	Brownerd
657	670	- var. macrorhizus	descript .
657 658	671	cinereus	mound .
659	672	tomentosus	No. American
		- niveus	sur crottin, non sur aiguille
66o	673	- micaceus	! sublime de l'art
661	674	- aratus	micaceus forme
662	675	- aratus	domesticus
663	675 676	— radians	Amening
			Psathyrella crenata
664	677	- alternatus	Panaeolus phalenarum
665	677 678	- deliquescens	male, exstinctorius?
666	719	- tardus	***************************************
667	679 680	congregatus	straight.
668	680	— Hendersonii	Automatic Control of the Control of
		- narcoticus	niveus
669	681	lagopus	-9/3/mmm
670	682	— macrocephalus	Psathyra pennata
100		- nycthemerus	malė
671-	683	radiatus	malè
200	-32	- Spraguei	Psathyrella crenata
672	684	— domesticus	malè
673	685	- stercorarius	malè
_	1	- ephemerus	teransia.
674	686	- plicatilis	malè
		- filiformis	sceptrum
675	687	- hemerobius	hiascens
0.0		- platypus	Advanta
676	688	Hiatula Wynniae	Coprinus narcoticus?
677	689	Bolbitius Boltoni	vitellinus
678	928	- vitellinus	Hygroph. chlorophanus
	100	- rivulosus	titubans ou Gal. tenera
679	720	— fragilis	titubans
	1 100	— apicalis	forme maladive
680	690	- titubans	

Maire	Rea	No. of bound volume	No. printed on Plat
orrugis?	gracilis	635 636	634
	hiascens	636	635
	· -	637 638	636
		638	655
Copr. plicatilis ou crenatus?			
audata	caudata, large form	639	637
lisseminata	disseminata	640	656
		-	× .
Vauc. temulenta		641	657
lisseminata	disseminata		
orrugis		642	638
gracilis	gracilis		
	crenata typical	643	847
omatus	comatus	644	658
ovatus	ovatus var. of comatus	645	659
terquilinus	sterquilinus	645 646	659 660
terquilinus forma	oblectus	647	661
utramentarius	atramentarius	647 648	662
itramentarius forma	atramentarius var. soboliferus	649	848
nicaceus, trop foncé	fuscescens	650	663
nicaceus ?	fuscescens var.	651	664
bicaceus	picaceus	652	665
Hyph. lacrimabundum Fr.		653	666
2	flocculosus	654	667
extinctorius		655	668
fimetarius	cinereus	656	669
fimetarius	macrorhizus	657 658	670
cinereus	cinereus	658	671
tomentosus		659	672
niveus	niveus		
micaceus	micaceus	660	673
micaceus?	-	661	674
?		662	675 676
radians	radians	663	676
?	Marine S. C.		100
?		664	677
?		665	678
Psathyra cernua?	Accident	1 666	719
congregatus	congregatus	667	719 679
Hendersonii	Hendersonii	668	680
5			1
lagopus	lagopus	669	681
?	1	670	682
?			
radiatus?	radiatus	671	683
crenatus?		1	
?		672	684
?		673	685
ephemerus	ephemerus		000
plicatilis	plicatilis	674	686
?	I		00
plicatilis	hemerobius	675	687
?	-	0.0	000
P		676	688
vitellinus?	vitellinus	677 678	689
vitellinus?	vitellinus	678	928
?	T * '		
	fragilis	679	720
titubans			
titubans apicalis	titubans	680	690

No. of bound volume	No. printed on Plate	Co	OKE	Quélet
68ı	691	Bolbitius tener	and the second s	Galera apala Fr.
682	692	Cortinarius (Phleon	nacium) triumphans	crocolitus
683	693		claricolor	Name of the last o
684	694		turmalis	claricolor
	695	Millionia Millionia	crussus	torvus, vetustior
685 686	696	Track there	balteatus	variicolor!
687	697	territoria 191.000.00	sebaceus	benè
687 688		*****	lustratus	argenteus
689	799 698	Marie	varius	ments.
690	699		cyanopus	Tips neare
691	700	Mary Access	variicolor	torvus
692	863	Name 1971	- var. nemorensis	Premise
693	701	menta Prome	largus	malė
694	702	tenteropy drawns	Riederi	Angelonia,
695	, ,	Ministry Account	saginus	triumphans forme
696	703	Orașional Windows	russus	brunneo-fulvus
697	751		infractus	cotoneus ou prasinus
698	704		anfractus	infractus
699	705 706		Berkeleyi	torvus! vetustus
			Berkeleyi	toreus
700 701	707 708	many many	multiformis	107 (16)
		many destroy	- var. flavescens	
702	709	Annana Minney	napus	turbinatus
703	710		allutus	Lactarius pubescens ou
704	752	memoral may be a second		Cort. cinnabarinus décolor
705	711	*****	talus	A 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
706	712	American Windows	glaucopus	Wis 4 dir M
707	713		calochrous	Transaction of the second
708	721	Constitute Productivity	coerulescens	pas la teinte
709	722	урцинц бирлор	coerulescens	{ dibaphus }
710	723	Manage Services	purpurascens	1
711	724	Windows Specialis	purpurascens	àti desiam
712.	725	-	- var. sub-	Personne
			purpurascens	
713	753	Marine Marine	dibaphus var.	turbinatus forme
0			xanthophyllus	
714	714	parties properly	turbinatus	Makeria
715	715	the same of the sa	corrosus	Solver -
716	716	emanag empleme	fulgens	Ange pair is
717	717	OTTO STATE OF THE	fulmineus	ne year
718	754	-	orichalceus	benè
719	735	Manual deposes	prasinus	Ser a more
720	736	- Accepted	atro-virens	Post Noted
721	755	-	scaurus	malè
722	849	-	herpeticus	glaucopus
723	726	The same of the sa	cumatilis	malė
724	727	Mining Property	emollitus	ochroleucus
725	728	Ministra - Process	cristallinus	eaustieus
726	729	-	decoloratus	newtood
727	730	-	decolorans	9.999 Ma
728	731	-	porphyropus	distance with the second secon
729	732	The same of the sa	croceo-coeruleus	malè
730	733	Manager Manager	corruscans	sebaceus
731	718		papulosus	multiformis
732	737 738	— (Myxacium)	arvinaceus	sebaceus
733		Transport American	collinitus	— (luxurians)
			mucosus	malè
734 735 736	739 740		mucifluus	collinitus

Maire	REA	No. of bound volume	No. printed on Plate	
Galera apala?	tener	681	691	
crocolitus	triumphans	682	692	
claricolor forme	claricolor	683	693	
turmalis		684	694	
praestans Cordier		685	695	
variicolor	balteatus	686	606	
	sebaceus	687	696	
argenteus?	Seouceus	688	697	
varius? plutôt forme de largus	varius		799 698	
largus jeune?	1	689	600	
praestans	cyanopus	690	699	
	praestans	691	700	
largus?	variicolor var. nemorensis	692	863	
largus	largus	693	701	
twitten to be an a	to invest to a	694	702	
triumphans	triumphans	695	703	
brunneo-fulvus?		696	75 ¹	
	infractus	697	704	
infractus	infractus	698	705	
praestans	praestans	699	706	
praestans	praestans	700	707	
multiformis	multiformis	701	708	
elegantior?	elegantior	702	709	
?	napus	703	710	
?		704	752	
multiformis	multiformis	705	711	
glaucopus	glaucopus	706	712	
calochrous	calochrous	707	713	
caerulescens var. caesio-cyaneus	caesio-cyanus	708	721	
caerulescens	caerulescens	709	722	
purpurascens	purpurascens	710	723	
purpurascens	purpurascens	711	724	
purpurascens var.	subpurpurascens	712	725	
xanthophyllus	dibaphus var. xanthophyllus	713	753	
turbinatus	turbinatus	714	714	
fulgens	fulgens	715 716	715	
fulmineus	fulmineus			
orichalceus	orichalceus	717	717	
prasinus	prasinus	, ,	754	
atro-virens	atro-virens	719	735	
2 ano-onens	scaurus	720	736	
glaucopus?	scaurus .	721	755	
giaucopus: cumatilis	cumatilis	722	849	
cumaturs emollitus	emollitus	723	726	
emotitus cristallinus		724	727	
	crystalinus decoloratus	725 726	1 .	
decoloratus	decolorans		729	
decolorans		727 728	730	
porphyropus	porphyropus croceo-caeruleus		731	
croceo-caeruleus	croteo-caermens	729	732	
5	papulosus	73° 731	733 718	
?		732		
collinitus	collinitus	733	737 738	
mucosus	mucosus	734	739	
collinitus	collinitus	735	740	

No. of bound volume	No. printed on Plate	Co	OKE	Quélet
737	742	Cortinarius (Myxae	cium) elatior	Biographic recognising and recognising development and property in a 1 of definition between the first and the fir
738			grallipes	sebaceus
739	734 767	grants According	livido-ochraceus	albo-cyaneus décoloré
740	768	-	salor	Monada
741	743		delibutus	eter-spot
742	831		stillatitius	diri-Mans
743	744		vibratilis	100.00446
744	769		pluvius	envisor.
	745	- (Inoloma)	argentatus	Antonoug
745 746	746	-	- var. pinetorum	amples
747	770	wateress and an artificial	violaceus	purpurascens
747 748	770 815	antoning months	muricinus	***************************************
749	747	privates privates	albo-violaceus	malè
750	756	-	malachius	glaucopus
751	771	promote desired	camphoratus	male, varius?
752	757	Name of the latest the	traganus	malè, amethystinus!
753	772	Military. Afficiacy.	tophaceus	-
754	773	-	redimitus	fulmineus
755	774		callisteus	percomis ou triumphans
755 756	774 864	months position	callisteus	hinnuleus
757	758		Bulliardi	malè
758		-	vinosus	purpurascens
759	759 760		bolaris	
759 760	761		pholideus	phylodologi
761	762	*****	sublanatus	(Miller)
762	763	-	arenatus	Mileson
763	764		penicillatus	Manage
764		- (Dermocybe	ochroleucus	persona,
765	775 816	(decumbens	causticus
, ,			diabolicus	rigens
766	783	-	tabularis	decoloratus
767	784	-	camurus	hinnuleus
767 768	765	Principle advanta '	caninus	delibutus
769	817		myrtillinus	benè!!
770	766		azureus	infractus
771	748		albo-cyaneus	sebaceus
772	776	-	anomalus	imbutus décoloré ou decolor
	0		1.612.4	tus
773	850		lepidopus	caninus
774	785	-	miltinus	weeks
775	786		cinnabarinus	man 1A
775 776		-	sanguineus	malè
110	787		anthracinus	human of all and
444	777		orellanus	brunneofulvus
777 778	777		cinnamomeus — var.	Normal Control of Cont
	770			danger.
779	779		- var. semi-	Nicerone
780	780		sanguineus	
100	,50	-	- var. croceus	
			conus	
781	851		uliginosus	
782	781		infucatus	* -
783	1 1		cotoneus	
784	749 832		subnotatus	raphanoides
784 785 786	750	-	valgus	
1-7	833		raphanoides	raphanoides venetus, vetustior
786			I MINIMANIUMEA	DEDELIK VEHISIOF
786	-33		venetus	- Contract Percention

Maire	REA	No. of bound volume	No. printed on Plate	
latior	elatior	737	742	
		738	734	
calor	livido-ochraceus	739	767	
	7 777	740	768	
llibatus (spore allongée)	delibutus typical	741	743	
ibratilis		742	831	
to antica	vibratilis	743	744	
urgentatus	pluvius typical	744	769	
rgentatus	argentatus	745	745	
ourpurascens?	argentatus var. pinetorum	746	746	
nuricinus	purpurascens	747 748	770 815	
argentatus?	albo-violaceus			
)	atoo-vioraceus	749	747	
	-	750	756	
raganus	twa a amus	75 T	771	
ophaceus	traganus	752	757	
2	tophaceus	753	772	
5		754	773	
innuleus	-	755 756	774 864	
Bulliardii ?	Bulliardii poor			
burpurascens?	vinosus	757	758	
polaris	bolaris	758	759 760	
pholideus	pholideus	759	761	
p	pholideus	760 761	762	
	pholideus	762		
penicillatus	photiacus	763	763 764	
ochroleucus	ochroleucus	764		
ristallinus?		765	775 816	
igens		/03	0.0	
decoloratus	——————————————————————————————————————	766	783	
ninnuleus?	-	767	784	
certe non delibutus; caninus? sporae	caninus poor	768	765	
differunt		/	7-5	
nyrtillinus	myrtillinus	769	817	
nfractus pâle?		770	766	
rebaceus?		771	748	
?	anomalus	772	776	
caninus	lepidopus	773	850	
phoeniceus (= miltinus Quél.)	phoeniceus	774	785	
cinnabarinus	cinnabarinus		-00	
plutôt la teinte de cinnabarinus	? cinnabarinus	775	786	
bhoeniceus?	anthracinus poor	776	787	
prellanus Fr. non Quél.	orellanus			
cinnamomeus	cinnamomeus	777	777	
cinnamomeus	cinnamomeus	778	778	
semisanguineus	semi-sanguineus	779	779	
cinnamomeus var. croceus	var. croceus	780	780	
cinnamomeus var. croceoconus	var. croceoconus	/55	1,00	
cinnamomeus var. uliginosus	uliginosus	781	851	
		782	781	
cotoneus	sublanatus	783	749	
raphanoides	subnotatus	784	832	
raphanoides	_	785	750	
venetus		786	833	
venetus	venetus	787	788	

No. of bound volume	No. printed on Plate		Co	OKE	Quélet
788	800	Cortinariu	s (Telam	onia) laniger	bivelus
789	852	*******	-	bivelus	more on
790	834	Mentury	****	bulbosus	trop rouge
791	818	******	traperiotis.	urbicus	- Telephone
792	819	pa-1-1-	WANT TOP	licinipes var . robustior	glandicolor
793	865		-	microcyclus	glandicolor
794	108		-	torvus	impennis!
795	853	ATTIONS	Ana.com	impennis	
795 796	820	-	*******	scutulatus	фтанци
		***************************************	Between	quadricolor	albo-cyaneus
797	821	******	*****	evernius	anomalus
797 798 -	866	******	#100 UM	evernius	elatior
799 800	867	delitore	and the same	quadricolor	elatior, gracilis
800	802	-	-	armillatus	haematochelis
108	803	-	Management	haematochelis	Who will de
802	804	-	alternative .	limonius	Tricholoma equestre?
	-	annou.	-	helvolus	hinnuleus
803	805	None of the last o	personal	hinnuleus	Search del
804	806	Permanen		gentilis	mal copié de Fries
80s	836	-	Married Ch.	helvelloïdes	obtusus var. gracitis
805 806	835	-	*****	rubellus	orellanus, major
807	822	*****	Mercen	bovinus	Winner
807 808	837	-	protection and the second	nitrosus	infractus, vetustior
809	823	Pro-	*****	injucundus	largus ou varius
810	854	******	Removal.	brunneus	impennis
811	854 868	Service in	SERVICE CO.	brunneus	Ministration
812	789	Things.	atomical .	glandicolor	milvinus?
_				· ·	(hinnuleus
813	855	-		punctatus	obtusus
814	790	-	-	triformis var. Schaefferi	claricolor
815	869	*****	*******	biformis	hinnuleus
816	838	*******	-	periscelis	copié de Fries
817	824	-	Minima.	flexipes	The same of the sa
/		PP - PR -	printerior in	flabellus	paleaceus (non vert)
818	839	-	Berneya .	psammocephalus	7
***	-33	-	Minimum Minimu	ileopodius	hinnuleus
819	807	***	-	incisus var. B.	leucopus?
820	825	******	-	hemitrichus	Tentopies .
821	840	Minney	intercomp.	stemmatus	hinnuleus
-	-1-	promise	-	Cookei	malè, spore fauve non jaun
822	791	*******	***************************************	rigidus	milvinus
823	826	-	Minney	paleaceus	majorial majorial
824		- (H	drocybe)	firmus	- Monora
825	792 808	President		subferrugineus	caninus
825 826		-	Percent	armeniacus	turinas
827	793 856	******	According	damascenus	2
828	827	***************************************	-	privignus	anomalus
829	809	**********		duracinus Fries?	armeniacus?
830	841	*******	Terrinos.	illuminus	with the state of
831	857		teritoria.	tortuosus	
832	810	-		dilutus	rigens
833	828		Moneyag	saturninus	*******
834	870	4		imbutus	impennis
835	842	-	-	castaneus	malè
836	871	1 10	-	bicolor	albo-cyaneus
	794				

glandicolor torvus Fr. non Quél. idem? scutulatus bicolor bicolor evernius? ? armillatus armillatus forma ? peut-être un Flammula ou Pholiota himuleus gentilis saniosus? ? ? largus? torvus Fr. non Quél.? brunneus ? potusus? } claricolor? ? flexipes? psammocephalus himuleus f. gracilis ?	bulbosus torvus impennis sscutulatus bicolor elatior quadricolor armillatus helvolus hinnuleus gentilis orellanus bovinus brunneus glandicolor	788 789 790 791 792 793 794 795 796 797 798 800 801 802 803 804 805 806 807 808 809 810 811 812	800 852 834 818 819 865 801 853 820 821 866 867 802 803 804 805 806 835 822 837 823 824 854 868
glandicolor torvus Fr. non Quél. idem? scutulatus bicolor bicolor evernius? ? armillatus forma ? peut-être un Flammula ou Pholiota himuleus himuleus gentilis saniosus? ? ? largus? torvus Fr. non Quél.? brunneus ? photusus? } claricolor? ? flexipes? psammocephalus himuleus f. gracilis ?	torvus impennis secutulatus bicolor elatior quadricolor armillatus armillatus helvolus hinnuleus gentilis — bovinus brunneus brunneus brunneus	789 790 791 792 793 794 795 796 797 798 799 800 801 802 803 804 805 806 807 808 809 810	852 834 818 819 865 801 853 820 821 866 867 802 803 804 805 806 835 822 837 823
glandicolor torvus Fr. non Quél. idem? scutulatus bicolor bicolor evernius? ? armillatus armillatus forma ? peut-être un Flammula ou Pholiota himuleus kinnuleus gentilis saniosus? ? ? largus? torvus Fr. non Quél.? brunneus ? flexipes? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	torvus impennis secutulatus bicolor elatior quadricolor armillatus armillatus helvolus hinnuleus gentilis — bovinus brunneus brunneus brunneus	790 791 792 793 794 795 796 797 798 799 800 801 802 803 804 805 806 807 808 809 810	834 819 865 801 853 820 821 866 867 802 803 804 805 806 835 822 823
glandicolor torvus Fr. non Quél. idem? scutulatus bicolor bicolor evernius? ? armillatus armillatus forma ? peut-être un Flammula ou Pholiota hinnuleus hinnuleus sinnuleus gentilis saniosus? ? ? largus? torvus Fr. non Quél.? brunneus ? pobtusus? claricolor? ? flexipes? psammocephalus hinnuleus f. gracilis ?	impennis scutulatus bicolor bicolor elatior quadricolor armillatus ————— helvolus hinnuleus gentilis —— orellanus bovinus —— brunneus brunneus	791 792 793 794 795 796 797 798 799 800 801 802 803 804 805 806 807 808 809 810	818 819 865 801 853 820 821 866 867 802 803 804 805 806 835 822 823
glandicolor torvus Fr. non Quél. idem? scutulatus bicolor bicolor evernius? ? armillatus armillatus forma ? peut-être un Flammula ou Pholiota hinnuleus hinnuleus gentilis saniosus? ? ? ? largus? torvus Fr. non Quél.? brunneus ? pobtusus? } claricolor? ? flexipes? psammocephalus hinnuleus f. gracilis ?	impennis scutulatus bicolor bicolor elatior quadricolor armillatus ————— helvolus hinnuleus gentilis —— orellanus bovinus —— brunneus brunneus	792 793 794 795 796 797 798 799 800 801 802 803 804 805 806 807 808 809 810	865 801 853 820 821 866 867 802 803 804 805 806 835 822 837 823
torvus Fr. non Quél. idem? scutulatus bicolor bicolor evernius? ? armillatus armillatus forma ? peut-être un Flammula ou Pholiota hinnuleus hinnuleus gentilis saniosus? ? ? largus? torvus Fr. non Quél.? brunneus ? potusus? } claricolor? ? flexipes? psammocephalus hinnuleus f. gracilis ?	impennis scutulatus bicolor bicolor elatior quadricolor armillatus ————— helvolus hinnuleus gentilis —— orellanus bovinus —— brunneus brunneus	794 795 796 797 798 799 800 801 802 803 804 805 806 807 808 809 810	801 853 820 821 866 867 802 803 804 805 806 835 822 837 823
idem? scutulatus bicolor bicolor evernius? ? armillatus armillatus forma ? peut-être un Flammula ou Pholiota hinnuleus hinnuleus sinnuleus gentilis saniosus? ? ? largus? torvus Fr. non Quél.? brunneus ? obtusus? claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	impennis scutulatus bicolor bicolor elatior quadricolor armillatus ————— helvolus hinnuleus gentilis —— orellanus bovinus —— brunneus brunneus	794 795 796 797 798 799 800 801 802 803 804 805 806 807 808 809 810	801 853 820 821 866 867 802 803 804 805 806 835 822 837 823
scutulatus bicolor bicolor evernius? ? armillatus armillatus forma ? peut-être un Flammula ou Pholiota hinnuleus hinnuleus gentilis ssaniosus? ? ? ? ? largus? torvus Fr. non Quél.? brunneus ? obtusus? } claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	scutulatus bicolor bicolor elatior quadricolor armillatus armillatus — helvolus hinnuleus gentilis — orellanus bovinus — brunneus brunneus	795 796 797 798 799 800 801 802 803 804 805 806 807 808 809 810	853 820 821 866 867 802 803 804 805 806 835 822 837 823
bicolor bicolor bicolor evernius? ? armillatus armillatus forma ? peut-être un Flammula ou Pholiota himuleus himuleus gentilis saniosus? ? ? ? largus? torvus Fr. non Quél.? brunneus ? pobtusus? } claricolor? ? flexipes? psammocephalus himuleus f. gracilis ?	bicolor bicolor elatior quadricolor armillatus armillatus helvolus hinnuleus gentilis — bovinus bovinus brunneus brunneus	797 798 799 800 801 802 803 804 805 806 807 808 809 810	821 866 867 802 803 804 805 806 836 835 822 837 823
bicolor evernius? ? armillatus armillatus forma ? peut-être un Flammula ou Pholiota hinnuleus gentilis saniosus? ? ? largus? torvus Fr. non Quél.? brunneus ? potusus? elaricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	bicolor elatior quadricolor armillatus armillatus helvolus hinnuleus gentilis orellanus bovinus brunneus brunneus brunneus	797 798 799 800 801 802 803 804 805 806 807 808 809 810	866 867 802 803 804 805 806 836 835 822 837 823
evernius? ? armillatus armillatus forma ? peut-être un Flammula ou Pholiota hinnuleus hinnuleus simuleus gentilis saniosus? ? ? largus? torvus Fr. non Quél.? brunneus ? obtusus? claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	elatior quadricolor armillatus armillatus helvolus hinnuleus gentilis orellanus bovinus brunneus brunneus brunneus	798 799 800 801 802 803 804 805 806 807 808 809 810	866 867 802 803 804 805 806 836 835 822 837 823
? armillatus armillatus forma ? peut-être un Flammula ou Pholiota hinnuleus hinnuleus gentilis saniosus? ? ? ? largus? torvus Fr. non Quél.? brunneus ? obtusus? claricolor? ? flexipes? psammocephalus hinnuleus f. gracilis ?	quadricolor armillatus armillatus ————————————————————————————————————	798 799 800 801 802 803 804 805 806 807 808 809 810	867 802 803 804 805 806 836 835 822 837 823
armillatus armillatus forma ? peut-être un Flammula ou Pholiota hinnuleus gentilis saniosus? ? ? largus? torvus Fr. non Quél.? brunneus ? obtusus? } claricolor? ? flexipes? psammocephalus hinnuleus f. gracilis ?	ormillatus armillatus helvolus hinnuleus gentilis — collanus bovinus — brunneus brunneus	799 800 801 802 803 804 805 806 807 808 809 810	802 803 804 805 806 836 835 822 837 823
armillatus forma ? peut-être un Flammula ou Pholiota hinnuleus gentilis saniosus? ? ? largus? torvus Fr. non Quél.? brunneus ? ? obtusus? claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	armillatus helvolus hinnuleus gentilis orellanus bovinus brunneus brunneus	801 802 803 804 805 806 807 808 809 810	803 804 805 806 836 835 822 837 823
? peut-être un Flammula ou Pholiota hinnuleus gentilis saniosus? ? ? ? ! largus? torvus Fr. non Quél.? brunneus ? obtusus? } claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	helvolus hinnuleus gentilis orellanus bovinus — brunneus brunneus	802 803 804 805 806 807 808 809 810	804 805 806 836 835 822 837 823
hinnuleus hinnuleus hinnuleus gentilis saniosus? ? ? largus? torvus Fr. non Quél.? brunneus ? obtusus? claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	hinnuleus gentilis — Orellanus bovinus — brunneus brunneus	803 804 805 806 807 808 809 810	804 805 806 836 835 822 837 823
hinnuleus gentilis saniosus? ? ? ? ! largus? torvus Fr. non Quél.? brunneus ? obtusus? obtusus? claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	hinnuleus gentilis — Orellanus bovinus — brunneus brunneus	804 805 806 807 808 809 810	806 836 835 822 837 823
gentilis saniosus? ? ? ? ? largus? torvus Fr. non Quél.? brunneus ? ? obtusus? claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	gentilis orellanus bovinus brunneus brunneus	804 805 806 807 808 809 810	806 836 835 822 837 823
saniosus? ? ? ? ? largus? torvus Fr. non Quél.? brunneus ? ? obtusus? claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	orellanus bovinus — brunneus brunneus	805 806 807 808 809 810 811	836 835 822 837 823
? ? ? ? ? largus? torvus Fr. non Quél.? brunneus ? ? obtusus? claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	bovinus 	806 807 808 809 810 811	835 822 837 823
? largus? torvus Fr. non Quél.? brunneus ? obtusus? claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	bovinus 	806 807 808 809 810 811	822 837 823
? largus? torvus Fr. non Quél.? brunneus ? ? obtusus? ? claricolor? ?	brunneus brunneus	808 809 810 811	837 823
torvus Fr. non Quél.? brunneus ? obtusus? obtusus? claricolor? flexipes? psammocephalus hinnuleus f. gracilis ?	brunneus	811 809	823
torvus Fr. non Quél.? brunneus ? obtusus? claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	brunneus	811	
brunneus ? ? obtusus? claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	brunneus	118	854 868
? ? obtusus ? } claricolor ? ?			868
? obtusus? } claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?	glandicolor	812	
claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?		1	789
claricolor? ? flexipes? ? psammocephalus hinnuleus f. gracilis ?		813	855
? flexipes? ? psammocephalus hinnuleus f. gracilis ?		814	790
flexipes? ? psammocephalus hinnuleus f. gracilis ?			
flexipes? ? psammocephalus hinnuleus f. gracilis ?		815	869
? psammocephalus hinnuleus f. gracilis ?	periscelis	816	838
hinnuleus f. gracilis	flexipes —	817	824
?	psammocephalus	818	839
	indiana.	0.0	907
?	incisus homituishaa	819	807
	hemitrichus	821	825
Cookei		1021	840
	rigidus	822	707
	paleaceus	823	791 826
pateateus — jienipis Ricken		824	
subferrugineus	subferrugineus	825	792 808
armeniacus	armeniacus	826	793
2	-	827	856
è		828	827
duracinus?	duracinus	829	809
?	<u> </u>	830	841
?	-	831	857
rigens?	Minimagene	832	810
saturninus	saturninus, pileus darker than usual	833	828
?	usuai	834	870
[castaneus]	castaneus, fairly typical	835	842
? figures inférieures		1	
bicolor balaustinus, trop rouge	bicolor typical	836	871

No. of bound volume	No. printed on Plate	Cooke	Quálet
838	795	Cortinarius (Hydrocybe) colus	obtusus, major
839	829	- isabellinus	
039	782	- renidens	anana.
840		- uraceus	Mineral II
841	796	jubarinus	hinnuleus
842	797 858	pateriformis var.	duracinus?
843	858	major	
844	859	- unimodus	erythrinus
845	811	- dolobratus	armeniacus!
845 846	812	- rigens	proper see
847	813	- Krombholzii	Mile with
847 848	843	- Reedii	milvinus
040	043	- leucopus	Non-see
0.0	000	scandens	leucofrus
849	830	- erythrinus	decipiens
850	798		hinnuleus
851	844	- germanus	malè
852	845	obtusus	yanning 1.5
		acutus	malè
853	846	— Junghuhnii	orellanus
	1	— milvinus	male, pas jaune
0	00-		duracinus
854	86o	— depressus	
855 856	814	- fasciatus	malè, stemmatus non jaur
856	879	Gomphidius glutinosus	Worked .
857 858	880	roseus	malè, plus rouge
858	188	viscidus	grunge
850	882	- maculatus (var.)	gallery tod
859 860	883	— gracilis	-
861	872	Paxillus (Lepista) lepista	glandonia.
862	874	- panaeolus	helomorphus
002	0/4	- orcelloides	mundulus
06.	0-0	- extenuatus	geotropus, forme
863	873 861	- lividus	Trich, cinerascens
864			
865	862	revolutus	H. subradiatus var. lacmus
866	884	paradoxus	distance
867 868	875	— (Tapinia) involutus	malè, trop jaune
868	929	- leptopus	trop jaune
869	876	- atrotomentosus	malè
870	877	crassus	vinosus Bull. = leptopus
877	877 878	panuoides	malè
871	0/0	Hygrophorus (Limacium) chrysodon	1 bené
872	885 886	rrygropnorus (Limacium) enrysouon	; Dene
873		- eburneus	
874	887	cossus	enservy
875	895	- pulverulentus	malè, non le mien
		- penarius	malè
876	888	- erubescens	
877	911	- pudorinus	prisoned
877 878	889	— glutinifer	olivaceo-albus, malè
879	896	- arbustivus	bearing .
-19	Ogo .	- aureus	Branch
880	912	- discoideus	malè, potius nitidus
881	897	Hygrophorus (Limacium) limacinus	malè, non jaune olive
882	890	- olivaceo-albus	malè, gracilis, vetustior
883	891	- hypothejus	Russula nauseosa
884	898	- cerasinus	Mangley
885	899	- fusco-albus	Lactarius trivialis

Maire	Rea	No. of bound volume	No. printed on Plate
5		838	795
?		839	795 829
renidens, trop pâle		840	782
uraceus	uraceus	841	796
?		842	
duracinus?		843	797 858
erythrinus	erythrinus	844	859
candelaris?	dolobratus	845	811
duracinus?	duracinus	846	812
?	Krombholzii	847	813
Inocybe sp.		848	843
leucopus	leucopus	040	043
p	Leavopas	849	830
erythrinus	and the same	049	
2 San thus	erythrinus	850	798
2	decipiens	0	0
14	germanus	851	844
obtusus	obtusus	852	845
acutus?	acutus		1
concinnus Karst. (= orellanus Quél.		853	846
non Fr.)			
?			
duracinus		854	860
?	fasciatus	855	814
glutinosus	viscidus typical	856	879
roseus	roseus	857	880
viscidus	viscidus	858	188
maculatus	maculatus	850	882
gracilis		859 860	883
gracuis	gracilis	861	
Ditti Trial-land 1. 1		862	872
Ripartites Tricholoma var. helomorpha	panaeolus typical	002	874
mundulus	orcelloides typical	00	0
Clitocybe amara Fr.		863	873 861
Clit. fumosa Fr. = cinerascens Quél.	lividus	864	
Hygr. pratensis var. cinereus? ou H. flavipes Britz. à cause des spores rondes		. 865	862
Phylloporus rhodoxanthus	paradoxus typical	866	884
involutus, trop jaune	involutus poor	867	875
non! spores differentes	- Poor	868	929
atrotomentosus, trop rouge	atrotomentosus	869	876
	ationientosus	870	877
? Phylloporus rhodoxanthus, vetustus	panuoïdes	871	878
panuoïdes		872	885
chrysodon	chrysodon		886
eburneus	eburneus	873	
eburneus var. cossus	cossus	874	887
niveus, forme à pied rosé?		875	895
penarius	penarius	1 0 0	000
erubescens	erubescens	876	888
pudorinus (forma fagetorum)	pudorinus	877	911
olivaceo-albus?	_	878	889
arbustivus	arbustivus	879	896
aureus	aureus		1
discoideus Fr., Lange = nitidus Quél. non Fr.	discoideus	880	912
olivaceo-albus?	- * *	188	897
olivaceo-albus, peu typique	olivaceo-albus, thin form	882	890
hypothejus, trop rougeâtre	hypothejus	883	891
cerasinus	cerasinus	884	898
fusco-albus?	fusco-albus	885	899

No. of bound volume	No. printed on Plate	Co	OKE	Quélet
886	913	Hygrophorus (Lime	ucium) agathosmus	male, gris, olivaceo-albus
887	914	20 1	mesotephrus	male, discoideus
888	915	- Indian	livido-albus	streptopus
889	916	(Camarophyl	lus) caprinus	7
890	930	armen armen	leporinus	
891	931	. series purious	nemoreus	Cortinarius armeniacus
892	917	personal statement	pratensis	ang or risk.
893	932	Section and section	- var. pallidus	malê
30	"		- var. cinereus	marc
894 895	892	yangkang Manang yantahing Affairing	virgineus — var. roseipes	niveus, major
		*******	niveus	gracilis
896	900	panents granten	russo-coriaceus	virgineus, minor
		part part	ventricosus	gliocyclus
897	901	anciently services	fornicatus	obrusseus, pâle
898	933	******	distans	clivalis
899	902	Market grant risk	Clarkii	waters
900	934	and the second	ovinus	malė
001	819	manage and and another a	metapodius	benè
901	935	-	subradiatus	representati
902	935	arrived dermany	- var. lacmus	MA LET
000	919	parameter alternated	irrigatus	and the second s
903	903	- (Hygrocyb	e) Colemannianus	spadiceus forme
904	903	()	foetens	Omphalia atropuncta, vieux
905	937	garage process	sciophanus	strature.
903	937		mucronellus	conicus, gracilis
906	938	·	laetus	* * **
907	936		Houghtoni	laetus! forme
908	904		vitellinus	Mirrogal
			ceraceus	t and and
909	920		coccineus	puniceus
910	921		miniatus	coccineus
			turundus var. mollis	Co. 1. 1. Liberta com viscon
911	905	-	Wynniae	Omphalia bibula var. viren
		anguage borners	micaceus	O. umbellifera var. flava
912	922	Accountage Notice and Accountage	puniceus	#AAND
913	906	Emergial School-	obrusseus	obrusseus
914	907	Reference participal p	intermedius conicus	ODI BISELIS
915	908	manuscript statement	calyptraeformis	
916	894	-	- var. niveus	amoenus var. alba
917	923		chlorophanus	coccineus décoloré
918	909		psittacinus	Barrer -
919	910		unguinosus	materia.
920	924		nitratus	caprinus pâle
921	925	Lactorius (Piher	tes) scrobiculatus	- Approximate the second secon
922	971	Zactartas (x tper	torminosus	malè
923 924	972 973	-	cilicioïdes	scrobiculatus, male
924	987	-	turpis	spin-maket
926	1003		controversus	(vetustus)
927	974	-	pubescens	torminosus (vetustior)
928	1083		aspideus	Stationer
929	975		insulsus	son and M
930	1084		utilis	Russula foetens?
931	988		blennius	
932	989		hysginus	malè
933	976		trivialis	zonarius (vetustus)
934	990		circellatus	

Maire	Rea	No. of bound volume	No. printed on Plate
? plutôt olivaceo-albus	71 17	886	913
leucophaeus (= discoideus Quél.non Fr.)	discoideus typical	887	914
Camarabhaller Fr - calminus	livido-albus	888	915
Camarophyllus Fr. = caprinus	camarophyllus	889	916
nemoreus, peu typique	leporinus	890	930
pratensis	pratensis	891	931
pratensis var. pallidus	pratensis pratensis var. pallidus	892 893	917
pratensis var. cinereus	pratensis var. cinereus	093	932
virgineus	virgineus	894	892
virgineus f. roseipes (= clivalis Q., teste Patouillard)	virgineus var. roseipes	895	893
virgineus var. niveus	niveus	896	900
virgineus var. russo-coriaceus	russo-coriaceus	3	
forme de virgineus		897	901
plutôt Trich. sejunctum grêle	clivalis	898	933
?	-	899	902
? .	irrigatus	900	934
ovinus	ovinus		
metapodius	metapodius	901	918
No. Anna pare	subradiatus	902	935
Post I am a de la companya del companya de la companya del companya de la company	lacmus		
irrigatus	irrigatus	903	919
Colemannianus	Colemannianus	904	903
O. atropuncta	foetens typical sciophanoides Rea	00"	007
sciophanoides Rea	mucronella	905	937
laetus	laetus	906	938
laetus	laetus	907	936
vitellinus	vitellinus	908	904
ceraceus	ceraceus	3	3-1
puniceus	coccineus, typical with yellow base	909	920
coccineus	miniatus	910	921
turundus var. mollis	turundus var. mollis	1.75	
O. Wynniae	Wynniae	911	905
pas O. umbellifera à cause des spores	micaceus		7.8
puniceus	puniceus	912	922
obrusseus	obrusseus	913	906
obrusseus	obrusseus	914	907
conicus calyptriformis	conicus calyptraeformis	915	894
calyptriformis var. niveus	calyptraeformis var. niveus	917	923
chlorophanus	chlorophanus	918	909
psittacinus	psittacinus	919	910
unguinosus	unguinosus	920	924
nitratus?	nitratus	921	925
scrobiculatus	scrobiculatus	922	971
torminosus	torminosus	923	972
?	cilicioides, pale form	924	973
turpis	turpis	925	987
controversus	controversus	926	1003
torminosus	pubescens	927	974
aspideus Fr. (flavidus Boud.)	uvidus	928	1083
insulsus	insulsus	929	975
Russ. delica?	blennius	930	1084
blennius	Ut and a second	931	
hysginus?	hysginus	932	989
insulsus, vieux?		933 934	990

No. of bound volume	No. printed on Plate	Co	OKE	Quélet
935	991	Lactarius (Piperites) uvidus	violascens!
936	992	trans.	flexuosus	uvidus (vetustus)
937	993	manual terminal	pyrogalus	or formation
938	1004	acresses a second	squalidus	
33-		Military Military	scoticus	argematus ou Russ, Raoultii
939	977	numbers distant	capsicum	rufus
940	984	Admin Town	chrysorrheus	december on
941	1005	Methodis. Paradish	acris	picinus Fr.
942	1006	Automa Marriago	umbrinus	Russula fusca Q.
943	978	property property	pargamenus	piperatus
944	979	manufig and comp	piperatus	and reported
	980	solution and advanced	vellereus	******
945 946	981	Annual recoded	exsuccus	Russula delica
	982	- (Dapetes) deliciosus	, sindrage
$\frac{947}{948}$	1007	(Russularia		ornor w
	983	A account to the contract of t	quietus	rufus
949 950	1099	transla busine	aurantiacus	Months and
	1008	anang mining	cremor var. pauper	tithymalinus?
951	1000	Berto ring	vietus	1014
952	1009	etionis branch	cyathula	non, argematus ou trivialis
050	1085	Manual Ma	cyathula	non! glyciosmus var. lilacinu.
953		Northean province	rufus	****
954	985	personal brokens	helvus	non! trop rouge
955	994	Manage McStreet	tomentosus	deliciosus on tithymalinus
956	1010	Topics or	mammosus var.	detroisas ou maymanas
957	995	market transport	monstrosus	
0		and his right	glyciosmus	2
958	1011	The state of the s		azonites ou picinus
959	996	protegue oursern	fuliginosus	azonaes ou pietnas
960	997	121 - 1	picinus	malè
961	998	- lilacinus		male
			s var. violaceus	man
962	999	— volemus — ichoratus		and a destar form of
963	1000	- icnoratus		espèce douteuse, forme d
964	1012	- serifluus		-000000
965	1001	- mitissimi	LS	official
966	1002	subdulcis		pas assez rouge
967	1013	- camphore	itus	subdulcis
3-1	3	- cimicariu		camphoratus
968	986	- subumbo	natus	subdulcis, forme approchar
				tabidus
		minimus		subdulcis var.
969	1014	- obnubilis	×	Hygr. hypothejus
		- obliquus		trivialis, forme
970	1015	- nigricans		adusta forma
971	1016	ls.	и	adusta, forme
972	1051	- adusta		adusta var. citrina
973	1017	- densifoli	1	nigricans?
974	1067	- semicrem		ochroleuca
	1068	- delica		var. albata
975 976	1018	- musteline	1	malè
077	1035	Russula olivascen		***************************************
977 978	1035	furcata	•	cyanoxantha, vetustior
970	1036		ar. pictipes	cyanoxantha, forme inconnu
979 980	1100		ar. ochroviridis	cyanoxantha, forme inconnu
981	1019	- sanguine)
982	1019	- rosacea	•	non, lepida
983	1020	- maculata		non! rien de commun; A
	1 1009	i iiiii iiiiiii		1 HORE LICH GC COMMUNE, P

Maire	Rea	No. of bound volume	No. printed on Plate	
uvidus	uvidus typical	935	991	
i'	huma a a la sa	936	992	
byrogalus ?	pyrogalus	937 938	993	
?		930	1004	
rufus?		939	977	
chrysorrheus	chrysorrheus	940	984	
picinus P		941	1005	
biperatus	piperatus var. pergamenus	942 943	978	
biperatus	piperatus typical	943	979	
vellereus	vellereus	945	980	
R. delica var. glaucophylla	R. chloroides	946	981	
deliciosus	deliciosus	947	982	
ballidus	pallidus	948	1007	
quietus, trop foncé aurantiacus	quietus	949	983	
état de volemus?	aurantiacus	950	1099	
vietus?	vietus typical	951	1008	
?	— typicar	952	1009	
lilacinus?		953	1085	
rufus	rufus	954	985	
paraît un vieux torminosus			994	
torminosus?		955 956	1010	
***************************************	-	957	995	
glyciosmus, peu typique	glyciosmus	958	1011	
bicinus	fuliginosus	959	996	
picinus	picinus	960	997	
lilacinus	lilacinus	961	998	
spinosulus var. violaceus	spinosulus var.			
volemus volemus	volemus	962 963	1000	
		3-3		
serifluus, trop violet	serifluus	964	1012	
mitissimus	mitissimus	965 966	1001	
quietus? peu typique camphoratus	ambharatus	966	1002	
camphoratus var. obnubilus	camphoratus cimicarius	967	1013	
camphoratus f. subumbonatus		968	986	
2			· ×	
: Hygr. hypothejus		969	1014	
?	× ×		_	
nigricans	nigricans	970	1015	
adusta var. albo-nigra	adusta var. albo-nigra	971	1016	
adusta	adusta	972	1051	
densifolia	densifolia	973	1017	
?		974	1067	
delica		975	1068	
mustelina?	mustelina	976	1018	
alutacea var. olivascens? cyanoxantha	alutacea var. olivascens	977 978	1035	
cyanoxantha forme	furcata var. pictipes		1086	
cyanoxantha, vieux	Jan out to the process of	979 980	1100	
sanguinea	sanguinea	981	1019	
sanguinea forme	sanguinea	982	1020	
depallens forme	atropurpurea	983	1069	

No. of bound volume	No. printed on Plate		Сооке	Quélet
984	1037	Russula	sardonia	non! decolorans ou olivascen var. citrina
985	1021		depallens	malė! amoena ou depallen var. vinosa
986	1022	montore.	purpurea	nitida ou palumbina, spore?
987	1052	******	coerulea	Management
988	1023	********	drimeia	Queletii
989	1070	-	lactea	manad
990	1071	Service of the latest	— var. incarnata	decembrate
991	1039	Marriagonia	virescens	non! pas vert olive
992	1024	-	cutefracta	- transition
993	1040		cutefracta	Min-state .
994	1072	APPRIATE	lepida	forme
995	1073	- Sussessed	lepida var.	white money
	'-		•	
996	1025	manipulari manipula m	rubra	sanguinea
997 998	1087	***************************************	rubra Fries (?) var. sapida Linnaei	reserved.
990 999	1026		xerampelina	? nitida
1000	1074	washed.	xerampelina	? inconnu
1001	1041		olivacea	
1001	1075	-	vesca	depallens var. vinosa ou
1003	1042	-	du Portii	cyanoxantha cyanoxantha
			serotina	Ar Hallo.
1004	1054	Market A	lilacea	Miles or many
1005	1088	where	azurea	All Andrews An
1006	1043	*******	cyanoxantha	bance
1007	1076	-	cyanoxantha	mercanica.
1008	1077	-	cyanoxantha var.	and the transfer of the state o
1009	1044	pa,ese	heterophylla	graminicolor
1010	1045	Minapag	heterophylla galochroa	foetens
1011	1089		consobrina	Promiti()
1012	1055		- var. intermedia	******
1013	1056		- var. sororia	49.00/01
1014	1057	Photos .	foetens	
1016	1047		subfoetens	? foetens, minor
1017	1058		fellea	Medit (g):
8101	1027		elegans	delica?
1019	1028	-	Queletii	- Manufacture
1020	1029	-	expallens	sanguinea, minor
1021	1030	Antiferencia	emetica	rosacea
1022	1031		Clusii	P
1023	1059	-	fallax	lilacea Q.
1024	1101	-	pectinata	foetens, minor
1025	1049	-	ochroleuca	
1026	1038	Processed .	granulosa	ochroleuca
1027	1090	-	aeruginea	- m- year
1028	1091		fragilis	Marin P
1029	1060	Heatenin	- var. violacea	aspect de lilacea
****	10.0	V	- var. niveus	Washing .
1030	1048	-	fingibilis	ochroleuca
1031	1078	**************************************	citrina Gillet	lutea
1032	1032	-	punctata var. leucopus	nitida forme

Maire	REA	No. of bound volume	No. printed on Plate
Queletii var. flavovirens?	-	984	1037
depallens = atropurpurea Krombh.	atropurpurea var. depallens	985	1021
depallens?	atropurpurea var. depallens	986	1022
caerulea	caerulea	987	1052
sardonia Fr. non Bres.	drimeia = sardonia Fr.	988	1023
lactea	lactea	989	1070
lactea var. incarnata	incarnata	990	1071
virescens)	virescens		1039
?		991	1
cutifracta cutifracta	cutifracta	992	1024
rosea Quél.)	cutifracta	993	1040
lepida }	lepida	994	1072
xerampelina jeune)	1.4.1.	1	
rosea Quél.	lepida	995	1073
atrorubens Quél.	atropurpurea typical	996	1025
melliolens forme foncée?	atropurpurea typical		1087
xerampelina var. erythropoda?	Linnaei typical	997 998	1026
grisea?	-	999	1053
xerampelina	xerampelina \	1000	1074
? fig. inférieure	fusca [
alutacea forme	olivacea typical	1001	1041
cyanoxantha forme?	vesca	1002	1075
xerampelina var.		1003	1042
serotina	serotina		1
vesca (sensu Bres.) ? ou lilacea géant	lilacea	1004	1054
azurea	azurea	1005	1054
cyanoxantha	cyanoxantha	1006	1043
idem .	cyanoxantha	1007	1076
grisea Bres.?	cyanoxantha	1008	1077
heterophylla	heterophylla	1009	1044
heterophylla forme	heterophylla	1010	1045
galochroa	galochroa	1011	1089
consobrina	consobrina	1012	1055
pectinata var.	consobrina var. sororia	1013	1056
idem	pectinata	1014	105
foetens	foetens	1015	1046
subfoetens sensu Maire=farinipes	laurocerasi	1016	104
Romell fellea	fellea	1017	105
elegans	Jetteu	1017	102
Queletii	Oueletii	1010	102
Queletii forme	drimeia	1019	102
rosacea Quél. non Fr.	emetica	1020	103
emetica	emetica var. Clusii	1022	103
fallax = olivaceoviolascens Gill.	emetica var. fallax	1023	105
pectinata?	pectinata	1024	110
ochroleuca	ochroleuca	1025	104
2	ochroleuca var. granulosa	1026	103
aeruginea? ou cyanoxantha forme	graminicolor	1027	109
fragilis	fragilis typical	1028	109
violacea Quél. ?	violacea	1029	106
fragilis var. nivea	fragilis var. nivea	1009	200
ochroleuca?	ochroleuca	1030	104
citrina Gill. non Quél.	citrina	1031	107
nauseosa Fr. forme = Turci Bres. sensu		1032	103
management I . I . I . I . I . I . I . I . I . I		1037	1 .03

No. of bound	No. printed	Cooke	Quélet
volume	on Plate		and the second s
1000	1033	Russula veternosa	up to some.
1033		- veternosa	chamaeleontina? spore?
1034	1092	- roseipes var.	xerampelina, gracilis, spore :
1035	1081		Attendation Brazilia, apara
1036	1034	integra	C
1037	1093	integra	fusca?
1038	1094	— var. alba	olivascens var. citrina, aspec
		11	de foctens
1039	1079	decolorans	2
1040	1061	Barlae	
1041	1080	aurata	2
1042	1062	— nitida var.	Barlae
1043	1063	nitida	rects and
		pulchralis	puellaris
1044	1095		fusca, minor
		— nitida var. cupraea	
1045	1064	armeniaca	chamaeleontina
1046	1065	- puellaris	With Park
1047	1066	- puellaris var.	nitida
1047	1096	- alutacea	Propries in
1048		alutacea	fusca?
1049	1097		Justin .
1050	1050	- ochracea	
1051	1082	lutea	Withday
1052	1147	nauseosa	Ministra b
1053	1102	— nauseosa var. flavida	lutea
1000	1102	- vitellina	ochracea
			at harms
		· ·	
1054	1098	chameleontina	au coin supérieur droit:
1055	1103	Cantharellus cibarius	Managem
1055		- cibarius var. rufipes	cibarius, vetustus -
1056	1131	- Friesii	cibarius
	-		
1057	1104		In sanish
1058	1106	- Brownii	albidus
		- umbonatus	Planter &
1050	1105	- carbonarius	- Man's midd
1059		- albidus	cibarius var. albescens
1000	1107	- Houghtoni	Marasmius incarnatus
			2 Managanta mentanta
1061	1108	- tubaeformis	
1062	1109	- infundibuliformis	
1063	1110	cinereus	cornucopiaides, forme
		- cupulatus	2.2
	16-	- leucophaeus	carbonarius, forme
1064	IIII	teucophaeus	
		- Stevensoni	albidus
1065	1115	- muscigenus	malè, il est cendré
		- glaucus	Borney.
1066	1112	retirugis	Month and
1000	1112	- lobatus	non, gris
C.			non, gris
1067	1132	Nyctalis caliginosa	parasitica
	. 1	- asterophora	# strang
1068	1113	- parasitica	N-MANA
1069	1116	Marasmius urens	paterings
		- peronatus	
1070	1117		Garidan 3
1071	1133	- porreus	foetidus?
1072	1118	- oreades	

Maire	Rea	No. of bound volume	No. printed on Plate
veternosa forme	veternosa	1033	1033
veternosa	veternosa var.	1034	1092
roseipes?	roseipes	1035	1801
Romellii Maire?	Romellii	1036	1034
integra Fr.?	integra	1037	1093
Romellii pâle?	Romellii	1038	1094
decolorans	decolorans	1039	1079
xerampelina var.?	xerampelina	1040	1061
aurata	aurata	1041	1080
?	-	1042	1062
nitida?	·	1043	1063
(nauseosa Fr.? puellaris?		1	
nitida	nitida var. pulchralis	1044	1095
	nitida typical		_
chamaeleontina	lutea var. armeniaca	1045	1064
puellaris	puellaris	1046	1065
xerampelina forma gracilis?	puellaris var. intensior	1047	1066
alutacea	alutacea	1048	1096
xerampelina	xerampelina	1049	1097
?	-	1050	1050
lutea	lutea	1051	1082
nauseosa Fr. (= Turci Bres.)		1052	1147
lutea? ou forme pâle de nauseosa		1053	1102
lutea forma? formes voisines de chamaeleontina: en haut à droite: lilacea Quél.	lutea var. vitellina,	-	
les 4 autres supérieures et les deux- inférieures: chrorosea Maire inédit; les 3 du milieu=R. abietina Peck	chamaeleontina	1054	1098
cibarius	cibarius	1055	1103
cibarius vieux	cibarius var. rufipes	1056	1131
cibarius			3
aurantiacus type et variété	\ \ aurantiaca \ \	1057	1104
	\ aurantiaca var. albida \	1	1
albidus		1058	1106
umbonatus	umbonatus		į
carbonarius	carbonarius	1059	1105
aurantiacus, forme décolorée		1060	1107
tubaeformis	tubaeformis	1061	1108
infundibuliformis	infundibuliformis	1062	1109
cornucopioides, forme à hymenium plissé	cinereus	1063	1110
	-		100
carbonarius		1064	IIII
albidus	-		
?	muscigenus	1065	1115
glaucus	glaucus		
retirugis	retirugis	1066	1112
?			
?		1067	1132
asterophora	asterophora poor	-	
parasitica	parasitica	1068	1113
peronatus, trop foncé	urens	1069	1116
peronatus	peronatus	1070	1117
?		1071	1133
oreades, trop jaune	oreades	1072	1118

No. of bound volume	No. printed on Plate	Сооке	Quélet
 1073	1119	Marasmius plancus scorteus	Collybia extuberans globularis
		prasiosmus	non, scorteus?
1074	1120	- varicosus	fusco-purpureus forme
1075	1121	fusco-purpureus	Mycena pura
		- terginus	an error
1076	1122	archyropus	globularis coloré
		Wynnei	globularis
1077	1123	erythropus	terginus
0	7704	- torquescens	Section Color
1078	1124	— impudicus	? malė
	1705	scorodonius	gazan similar
1079	1125	calopus	Vaillantii
1080	1126	Vaillantii	white
1000	1120	- angulatus	Vaillantii
		- languidus	Omph. umbellifera?
0	*****	foetidus	allestrade of
1801	1134	cauticinalis	Newmond
	1		
. 0 -		amadelphus	yang darah
1082	1127	- candidus	ggrade rol
		ramealis	MARKET MARKET
- 0 -	0	alliaceus	\$400.000A
1083	1128	cohaerens	fusco-purpureus
		- rotula	******
1084	1129	graminum	B-028999
		gramera	
		- androsaceus	?
		splachnoides	androsaceus
1085	1130	- Curreyi	graminum
		- perforans	malė
.00		- insititius	malė
1086	1135	- Hudsoni	gelidus Q.?
	16	epichloë	caulicinalis
1087	1136	- actinophorus	graminum
		- saccharinus	- Sept. of
- 00		- epiphyllus	papers.
1088	1137	polyadelphus	saccharinus
		- spodoleucus	Pleur. cyphellaeformis
0 -	0	Lentinus tigrinus	Amenito
1089		- tigrinus	No. Ashina
1090	1139	Dunalii	1,0000
8		- lepideus	distribution and
1091		— monstrous form	u ryphe
1092	1141	- Industrous form	
		- cochleatus	National
1093	1142	- vulpinus	potius flabelliformis Schaeff.
		1	suaveolens
1094		Scotteus	America .
1095	1148	- fimbriatus	No. of the last of
		— flabelliformis Panus conchatus	flabelliformis
1096	1149		flabelliformis
		- torulosus	January Comments of the Commen
109,	7 1144	- stypticus	flabelliformis forme d'hiver
		— farinaceus	3
		patellaris	inconnu
109	8 1150	Cantharellus devexus Xerotus degener	malè
		1 A FTO CITS TIE DETIET	
109	9 1114		*****

Maire	Rea	No. of bound volume	No. printed on Plate	
?	james and the same	1073	1119	
Wynnei, petite forme pâle				
fusco-purpureus?	varicosus	1074	1120 1121	
Mycena pura	- varicosus	1075	1121	
terginus		1076	1122	
?				
Wynnei	globularis old	1077	1123	
lupuletorum	erythropus typical	0		
torquescens impudicus	torquescens impudicus	1078	1124	
scorodonius		1079	1125	
Vaillantii forma		10/3	1	
Vaillantii	Vaillantii	1080	1126	
Vaillantii forma				
?		0-		
foetidus	foetidus poor	1081	1134	
fulvobulbillosus R. Fries, chapeau trop foncé	cauticinalis typical			
amadelphus	manufa,	1082	1127	
candidus	candidus			
ramealis	ramealis			
alliaceus	alliaceus	1083	1128	
?		0.		
rotula	rotula	1084	1129	
graminum?	graminum, stem should be bright			
androsaceus	androsaceus			
splachnoides		1085	1130	
graminum?				
perforans? trop pâle	perforans	06		
7	insititius Hudsonii	1086	1135	
Hudsonii stipitarius?	Huasomi	1087	1136	
supuarus :		1.557	1 3-	
saccharinus	saccharinus			
? spores trop petites	epiphyllus	1088	1137	
Omphalia polyadelpha	polyadelphus	,	100	
Pleur. cyphellaeformis	d'antique d'	1089	1138	
tigrinus	tigrinus tigrinus	1009	1139	
tigrinus tigrinus	igi mus	1090	1.39	
lepideus	lepideus	1091	1140	
lepideus, anomalie	lepideus, form not	1092	1141	
	uncommon			
cochleatus, trop rouge	cochleatus	1093	1142	
Panus rudis?	scoticus	1094	1143	
f Haminus	fimbriatus	1094	1148	
tigrinus torulosus?	Jimortanas	1.033		
torulosus	torulosus	1096	1149	
torulosus	torulosus			
stipticus	stypticus	1097	1144	
?				
3	yer of replement	1098	1150	
?	var. of replexus	1090	1130	
rispa	crispa	1099	1114	
commune	commune		1	

No. of bound volume	No. printed on Plate		Coor	KE .	QUÉLET
1100	1145	Lenzites betu		4	New york or
	10	flacc			460 (190)
1011	1146	- sepi	aria		erent of
		- abie	tina		saepiaria
1102	939	Agaricus (Ar	nanita)	solitarius	solitaria Quél.
1103	1163	-	Majoritorio	rubescens	Brong pr
1104	940	arran.		vaginatus var. nivalis	pas de violet
1105	941	(<i>l</i>	.epiota)	Friesii	malè
1106	1164	Afficia	Particolo.	emplastrum	Am, rubens! spore anormale, ou forme anormale de L. rachodes
1107	1180		Market .	hispidus	rating e.g.
8011	943	Personal	apertoresis	felinus	Month of
1100	943	******	-	micropholis	felina, gracilis
****	0.40	- American	W/10 W 100	cepaestipes var.	benê
1109	942			cretaceus	
	*****	-	garage.	liemothorus	cepaestipes var. lutea
1110	1179	7	access,	citrophyllus	cristata malade, ou clypeo-
IIII	639			can apagamen	laria, gracilis
	0.44		an broken	ianthinus	cristata, gracilis
1112	944			martialis	haematosperma, gracilis
		(12	millaria	Jasonis	Lep. amianthina
1113	955	(217)	munitin m	focalis var . goliathus	Lep. ampla P.
1114	1165			Citri	mellea, variété
1115	1181	177:	abalama'	russula	water the state of
1116	926	(111	cnowna	variegatus	******
1117	642			argyraceus var.	
1118	64.1		-	virescens	
			-	argyraceus var.	
1119	947			chrysites	•
1120	945		processa	inodermeus	Hygrophorus russula
	60			tenuiceps	cartilagineum on molybdinum
1121	1166	-	AN INTERNAL	fallax	album on inamoenum
1122	1151	- (4	Wite who) simararane	expallens
		- (6	attocyoe) cinerascens	expanens
- 25.00	6	17	ichaloma	borealis	Entoloma clypeatum
1123	956	- (11		pes-caprae	aggregatum
1124	946			circumtectus	argyraceum? ou luridum
1125	1182			duracinus	cinerascens
1126	640			melaleucus var.	grammopodium, minus?
1127	957			polioleucus	grammopourtant, treaters
1128	1183	_ (Clitocybo	e) opiparus	Trich. truncatum Sch., très bonne figure
1129	644		7	amplus	www.co
1130	645		-	fumosus	Name of the second
1131		_	-	subdecastes	Hygr. melizeus? on eossus vieux
1132	643			pergamenus	lignatilis
1132			***************************************	occultus	argyraceum, ou virgatum ou hordum
	648			monstrosus	Hygr, virgineus?
1134	1 0 0			infundibuliformis	JEI CHEMENS
1135	646	1 1 21 2		var. membranaceu	15
+ + = 0	64-		10	sinopicus	and the same of th
1136				zygophyllus a) fodiens	amarella

Maire	REA	No. of bound volume	No. printed on Plate	
betulina	betulina	1100	1145	
flaccida	flaccida		10	
saepiaria	saepiaria	1101	1146	
saepiaria, vieux	abietina		1	
echinocephala	echinocephala	1102	939	
rubescens	rubescens	1103	1163	
vaginata var. nivalis	nivalis	1104	940	
Friesii	Friesii	1105	941	
rhacodes	rhacodes	1106	1164	
hispida	hispida	1107	1180	
felina	felina, large form	8011	943	
7	micropholis		343	
cepaestipes var. cretaceus	var. cretaceus	1109	942	
	7 7			
lutea	licmophora	1110	1179	
fig. inf. citrophylla	citrophylla	IIII	639	
lilacea Bres.?		1112	944	
?				
amianthina	amianthina	1113	955 1165	
Amanita spissa forme	***	1114	1105	
mellea forme	mellea var.	1115	1181	
Hygrophorus Russula	Hygr. Russula	1116	926	
-	variegatum	1117	642	
scalpturatum forma	argyraceum var.	1118	641	
idem	argyraceum var.	1119	947	
Hygr. Russula? ou forme stérile de Inocybe piriodora	inodermeum, a good rare species	1120	945	
Coll. platyphylla forma?	aggregatum	1121	1166	
2		1122	1151	
P	cyathiformis, from stem			
	characters		0-6	
ľ		1123	956	
pas aggregata à cause des spores		1124	946	
atrosquamosum forma?	· Dull man	1125	1182	
cinerascens Quél. non Fr.	cinerascens Bull., poor	1126	640	
humilis forma	melaleucum var. polioleucum	1127	957	
truncatum	- *	1128	1183	
transformis Britz., si les spores sont triangulaires	, - - , _ +	1129	644	
aggregata var.	cinerascens	1130	645	
idem	cinerascens	1131	958	
idam	pergamena	1132	643	
idem ?	Russula? sp.	1133	1184	
•	Trasaua; sp.	1133	1104	
assuments forms ?	monstrosa	1134	648	
cerussata forma? infundibuliformis forma membranacea	infundibuliformis var. mem- branacea	1135	646	
singhiag	sinopica, small form	1136	647	
sinopica inornata	zygophylla	1137	948	
	fodiens	1138	340	

No. of bound volume	No. printed on Plate	Coc	DKE	Quélet
1139	950	Agaricus (Collybia)	prolixus	distorta
1140	652	province directors	distortus var.	Manager
1141	650	Minoritian Anno ya	velutipes var. rubescens	*** **********************************
1142	1168	bronung debroom	floccipes	inconnu
1143	1167	are obtained for the first	thelephorus	Mycena plicosa?
1144	651	decorage and house	leucomyosotis	Mycena rugosa
1145	649	Allegang Mil John	tenacellus	Fills !
1146	1185	state of the state	eustygius	Clit, inornata
1147	1198	THE STATE OF THE S	murinus	malè
1148	951	- (Mycena)	mirabilis	iris
	33	Marriage Marriage	flavipes	Militages and
1149	952	Managed Statements	gypseus	# Grap. pgs
		monage Windows	codoniceps	trachelina Fr.
1150	1186	breedly Arthury	consimilis	leptocephala
1151	653	Defense on Minutes	leucogalus	galopus
1152	1152	(Omphalia)) chrysophyllus	Mericon
		Service Service	Postii	-
1153	959	- (Mycena)) olivaceo-marginata	aurantio-marginata
		- (Omphalia) glaucophyllus	application
		******	rusticus	fings a
1154	654	- (Pleurotus) Ruthae	conchatus ou palmatus?
1155	954	Manage Marriage	sapidus	conchatus
1156	953	Military Military	columbinus	ostreatus
* 1157	1169) salicinus	malè
	-	- (Leptonia) asprellus	malè, il est gris bistre
1158	1153	- (Entoloma) nigro-cinnamomeus	inconnu
1159	960	- (Clitopilus) straminipes	Entoloma speculum
1160	1170	(Nolanea) nigripes	Naucoria cucumis
		/ F32 - 2* 1 ·	subglobosus	icterina
1161	1171) molliscorium	ombrophila
1162	961	- (Inocybe) perlatus	fastigiata, forme obèse
1163	1174	Jackson Sections	violaceo-fuscus	obscura
1164	1173	- (Hebeloma	fasciatus	pyriodora
1165 1166	962 963	- (Treveroma	nauseosus	longicandum
1167	964	- (Flanmula) purpuratus	Ph. confragosa? ou mieux
	904	(a tammata		Tr. variegatum?
1168	1154	protecting delicitation	nitens	Tr. phoenix Mich. ou pertina Fr. inconnu mais semble l
**60	770-	/ Massar-i	Lumbric	même espèce
1169	1187	(Naucoria	j tuguoris festivus	malè
1170	966		obtusus	Psil. sarcocephala Fr.
1172			hamadryas	inconnu
1173	965		nasutus	myosotis, variété
/3		(Pholiota) blattarius	Lep. echinata, décolore
7.1774	1156	- (Galera) siligineus	ou Heb. erebium Fr. gracil
1174	1175) pellucidus	apala, tenera, etc. aguosa?
11/3	11/3	(I abarta	muscorum	Galera Q.
1176	967	- (Chitonia) rubriceps	inconnu, espèce exotique
	968	- (Psaliota) sagatus	semota Fr., variété adulte
1177	1188	/a preference) merdarius var.	squamosa
1179	1189		major scobinaceus	Hubb abbondinglation
** 19	1109		scoothace as	Hyph, appendiculatum

Maire	Rea	No. of bound volume	No. printed on Plate
distorta?	***************************************	1139	950
listorta	distorta var.	1140	652
velutipes forma rubescens	velutipes var. rubescens	1141	650
Mycena maura Maire? trop grand		1140	1168
Mycena		1142	1167
3	laucomoncotic turnical	1143	
tenacella	leucomyosotis typical tenacella	1144	651
	ienacetta	1145	649
Trich. immundum Berk. = Ag. fumosus Pers. non Fr.		1146	1185
?	-	1147	1198
Iris forma?	marginella	1148	951
flavipes = Renati	flavipes	*	
yypsea?	gypsea	1149	952
?*	-	13	33
?		1150	1186
galopoda var. leucogala	galopus var. nigra	1151	653
chrysophylla	2	1152	1152
Postii		1132	1152
avenacea forme	avenacea var. olivaceo-	1750	050
avenuteu formic		1153	959
· ·	marginata		
	glaucophylla		
rustica	rustica		C
palmatus	palmatus	1154	654
cornucopiae	sapidus	1155	954
ostreatus var. columbinus	ostreatus var. columbinus		953
salicinus	salicinus	1157	1169
asprella	asprella	0	N. C.
?		1158	1153
?	Lept. sericella	1159	960
Nauc. Cucumis forme	Nauc. Cucumis	1160	1170
? e sporis potius Pluteus			
?	molliscorium	1161	1171
fastigiata forme	perlata	1162	961
obscura	violaceofusca	1163	1174
3	fasciata	1164	1173
elatum	elatum	1165	962
? ressemble à <i>H. sacchariolens</i> mais		1166	963
spore trop grande		1167	064
r	,	1107	964
? voir formes brunes exannulées de Ph. cylindracea		1168	1154
lugubris	lugubris	1169	1187
festiva	festiva	1170	966
)		1171	1155
5		1172	965
Psil. semi-lanceata		1173	1172
?	blattaria	1.75	1
apala et tenera?	Galera campanulata	1174	1156
para de tenera.		1175	1175
•		,3	,,,
rubriceps	rubriceps	1176	967
rubella Gill?		1177	968
stercoraria forma? ou plutôt Hyph.		1178	1188
capnoides	*		0
Hyph. lacrimabundum Fr.? ou forme	scobinaceum	1179	1189

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1180	1176	Agaricus (Hypholoma) catarius	Hyph, appendiculatum
1811	1157	- instratus	hydrophilum
1182	1177	- (Psilocybe) areolatus	Stroph, Battarrae?
1183	969	- Clivensis	Heb. elatum gracile ou Heb. sacchariolens
1184	970	- (Psathyra) gyroflexus	Page 11
1185	1158	conopileus var . superbus	Psathyra fatua
1186	1160	- (Bolbitius) conocephalus	vitellinus, blanchi; ou apalus exubérant
1187	1159	grandiusculus	vitellinus var. Boltonii
1188	1190	Cortinarius (Phlegmacium) testaceus	subpurpurascens Fr.
1189	1191	(Myxacium) nitidus	sebaceus forma
1190	1192	- (Telamonia) lucorum	non, evernius
1191	1193	croceo-fulvus	limonius Fr.
1192	1178	— (Hydrocybe) angulosus	cinnamomeus forma
1193	1162	Paxillus Alexandri	malė
1194	1161	Hygrophorus (Hydrocybe) spadiceus	1 40 4 11
1195	1194	Lactarius involutus	piperatus ou argematus?
1196	1195	squalidus	inconnu: blemius, vietus?
1197	1197	Russula virginea	delica, forme on lactea
1198	1196	— ochroleuca var. claroflava	olivascens var. citrina

Maire	Rea	No. of bound volume	No. printed on Plate	
Candolleanum forma hydrophilum? spore trop grande Hyph. melantinum ?	catarium clivensis	1180 1181 1182 1183	1176 1157 1177 969	
? Psathyrella subatrata	gyroflexa	1184 1185	970 1158	
?	- *	1186	1160	
vitellinus var.? rufo-olivaceus, adulte, très probable- ment	testaceus	1187 1188	1159	
? torvus Fr. non Quél.?	_	1189	1191	
tophaceus Fr. Ricken! venetus?	* ×	1191	1193 1178 1162	
spadiceus D'après les spores ce serait plutôt		1193 1194 1195	1161	
un Clitocybe du groupe cerussata blennius?	-	1196	1195	
? forme blanche de quelque autre espèce flava Romell ? et forme de R. ochroleuca		1197	1197	

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369 378 417 466 475 476 474 534 557 592 610 649 1145 360 381 418 461 476 476 474 534 557 592 612 650 1141 361 382 419 462 477 482 533 558 593 615 651 1144 362 383 420 463 478 502 535 558 593 615 651 1144 362 383 420 463 478 502 535 558 593 615 651 1144 363 384 421 313 479 506 537 565 595 618 653 1151 364 385 422 315 480 513 538 566 596 622 654 1154 365 387 423 380 481 521 539 567 597 308 655 638 366 389 424 406 482 510 540 568 598 312 656 640 367 391 425 409 483 528 541 569 599 350 657 641 368 396 426 419 484 529 542 570 600 388 658 644 369 397 427 421 485 346 543 580 601 495 659 643 370 398 428 435 486 347 544 584 602 526 660 646 370 391 428 429 429 439 487 351 545 585 603 527 661 647 372 403 430 448 488 360 546 586 604 564 662 646 373 373 323 431 465 489 498 547 587 605 589 663 557 374 336 432 476 490 500 548 588 606 599 664 665 648 375 345 433 477 491 501 549 586 600 666 653 376 362 434 478 492 505 550 554 608 600 666 653 377 363 435 479 493 507 551 555 609 601 6667 668 380 380 448 440 470 498 535 556 553 600 601 6667 663 380 380 444 444 481 502 399 560 575 613 600 666 663 638 380 448 440 470 498 535 556 563 611 614 669 654 380 431 447 439 499 497 533 555 566 610 606 688 633 380 448 449 490 500 548 588 600 691 6672 6639 380 441 449 490 500 548 588 573 610 611 614 669 656 383 384 418 440 470 498 535 556 563 611 614 669 665 380 441 443 480 501 348 559 574 661 612 621 621 670 657 380 443 445 499 501 534 557 572 613 648 663 673 660 386 437 446 481 502 399 560 575 618 553 669 611 667 663 380 441 444 481 502 399 560 575 618 553 669 611 677 664 381 447 449 490 507 447 487 509 496 567 575 618 553 669 611 674 661 672 621 621 621 621 621 621 621 621 621 62	357		415	450		471				608	648	
36io 38i 418 46i 476 474 534 557 592 612 651 6114 36i 38g 419 462 477 48g 535 558 593 615 651 1144 36i 38g 420 463 478 502 536 559 594 616 652 1140 36i 38j 422 315 480 513 538 566 596 622 654 1154 36i 38g 424 406 482 510 540 568 598 312 656 640 36i 38g 424 406 482 510 540 568 598 312 656 640 36i 38g 424 409 483 528 541 569 599 350 657 641 36i 39i 427 421 485 346	359	378	417	460		472				610	649	1145
362 383 420 463 478 502 536 559 594 616 653 11451 363 384 421 313 479 506 537 565 359 594 616 653 1151 364 385 422 315 480 513 538 566 396 622 654 1154 365 387 423 380 481 521 539 567 597 308 655 630 366 389 424 406 482 510 540 568 598 312 656 640 367 391 425 409 483 528 541 569 599 350 657 641 368 396 426 419 484 529 542 570 600 388 658 644 369 397 427 421 485 346 543 580 601 495 659 645 370 398 428 435 486 347 544 584 602 526 660 646 331 400 429 439 488 350 546 586 604 564 662 643 373 322 403 448 488 360 546 586 604 564 662 643 373 323 431 405 499 500 548 588 606 599 664 651 375 376 362 434 478 492 505 550 554 608 601 667 654 377 363 435 479 493 507 551 555 609 601 667 654 378 379 366 437 466 495 516 553 560 611 614 669 656 380 368 438 440 470 498 532 554 561 612 621 670 653 381 417 439 449 499 532 554 551 652 613 369 671 653 382 418 440 470 498 532 554 561 612 621 670 653 383 426 441 473 499 534 557 572 613 369 671 653 384 448 440 470 498 535 556 613 369 671 653 384 449 442 475 500 488 557 572 613 369 671 653 384 440 470 498 532 554 561 612 621 670 657 381 447 439 449 499 534 557 572 613 369 671 653 382 418 440 470 498 535 556 563 614 489 674 661 384 449 442 475 500 488 559 574 617 504 663 386 431 444 481 502 399 560 575 618 553 666 663 387 443 443 488 501 348 559 574 617 504 661 384 449 442 475 500 498 535 556 613 369 671 653 384 440 470 498 535 556 563 614 393 674 661 385 430 443 480 501 348 559 574 617 504 675 662 386 437 446 484 504 470 508 575 575 618 553 666 663 673 660 666 386 437 446 484 504 470 508 552 556 619 606 666 668 655 673 660 666 668 655 673 660 666 668 655 673 660 666 668 655 673 660 666 668 665 673 674 660 660 666 668 665 673 674 675 675 675 675 677 660 663 669 677 677 664 677 677 664 677 677 664 677 677	360	381		461	476	474	534	557			650	
363 384 421 315 479 506 537 505 595 618 653 1151 364 385 422 315 480 513 538 566 556 622 654 1154 365 387 423 380 481 521 539 567 597 308 655 638 366 389 424 406 482 510 540 568 598 312 656 620 367 391 425 409 483 528 541 569 599 350 657 641 368 396 426 419 484 529 542 570 600 388 658 644 370 398 428 435 446 347 544 584 602 556 660 642 371 400 429 439 498 548 <t< td=""><td>361</td><td></td><td></td><td></td><td>477</td><td></td><td>535</td><td>550</td><td></td><td></td><td>652</td><td></td></t<>	361				477		535	550			652	
365 387 423 380 481 521 539 567 597 308 655 638 366 389 424 406 482 510 540 568 598 312 656 640 367 391 425 409 483 528 541 569 599 350 657 641 368 396 426 419 484 529 542 570 600 388 658 644 369 397 427 421 484 529 542 570 600 388 658 644 370 398 428 435 486 347 544 584 602 526 660 660 640 371 400 429 439 487 351 545 586 602 562 662 642 373 336 432 476 499	363				479	506	537	565			653	1151
365 387 423 380 481 521 539 567 597 308 305 636 640 367 391 425 409 483 528 541 569 599 350 657 641 368 396 426 419 484 529 542 570 600 388 658 644 369 397 427 421 485 346 543 580 601 495 659 645 660 644 369 397 427 421 485 346 543 580 601 495 659 645 660 644 371 400 429 439 487 351 545 584 602 526 660 640 642 432 433 431 405 489 498 547 587 605 589 663 650 663 632 373 336 432 477 491 <td></td> <td>385</td> <td></td> <td>215</td> <td>480</td> <td>513</td> <td>538</td> <td>566</td> <td>596</td> <td></td> <td>654</td> <td>1154</td>		385		215	480	513	538	566	596		654	1154
367 391 425 409 483 528 541 569 599 350 657 641 368 396 426 419 484 529 542 570 600 388 658 644 370 398 428 435 486 347 544 584 602 526 660 645 371 400 429 439 487 351 545 585 603 527 661 647 372 403 448 488 360 546 586 604 564 662 648 373 323 431 465 489 498 547 587 605 589 665 652 663 650 374 336 435 479 491 501 549 358 607 598 666 652 377 362 434 479 493				380	481			567	597	308	655	640
369 397 427 421 485 346 543 580 601 495 569 645 370 398 428 435 486 347 544 584 602 526 660 645 371 403 430 448 488 360 546 586 604 564 662 648 373 323 431 465 489 498 547 587 605 589 663 653 374 336 432 476 490 500 548 588 606 599 664 651 375 345 433 477 491 501 549 358 606 599 664 651 376 362 434 478 492 505 550 554 608 600 666 663 632 377 363 435 479 493			424		482			560	590	350	657	
369 397 427 421 485 346 543 580 601 495 569 645 370 398 428 435 486 347 544 584 602 526 660 645 371 403 430 448 488 360 546 586 604 564 662 648 373 323 431 465 489 498 547 587 605 589 663 653 374 336 432 476 490 500 548 588 606 599 664 651 375 345 433 477 491 501 549 358 606 599 664 651 376 362 434 478 492 505 550 554 608 600 666 663 632 377 363 435 479 493			426		484	529		570	600	388	658	644
371 400 429 439 487 351 545 585 603 527 661 647 372 403 439 448 488 360 546 586 604 564 662 648 373 323 431 465 489 498 547 587 665 589 664 651 374 336 432 476 490 500 548 588 600 599 664 651 375 345 433 477 491 501 549 358 607 598 665 652 376 362 434 478 492 505 550 554 608 600 666 653 377 363 435 479 493 507 551 555 609 601 667 654 378 363 435 479 493 507 551 555 609 601 667 654 378 363 435 479 493 507 551 555 609 601 666 658 379 366 437 466 495 516 553 560 611 614 669 656 380 368 438 467 496 532 554 561 612 621 670 657 381 417 439 469 497 533 555 562 613 369 671 658 382 418 440 470 499 534 557 572 615 486 673 660 384 442 442 475 500 468 558 573 616 489 674 661 385 430 443 480 501 348 559 574 617 504 675 662 386 431 444 481 502 399 560 575 618 553 676 663 387 433 445 488 501 348 559 574 617 504 675 662 389 407 447 487 505 432 563 582 577 620 571 677 664 388 437 446 484 504 423 562 577 620 591 678 665 389 407 447 487 505 432 563 582 577 620 591 678 665 389 407 447 487 505 432 563 582 577 620 591 678 665 399 410 448 488 506 454 564 581 582 591 678 662 393 416 451 492 509 496 567 591 678 662 393 416 451 492 509 496 567 590 625 626 683 671 393 416 451 492 509 496 567 590 625 626 683 671 393 416 451 492 509 496 567 590 625 626 683 671 393 416 451 492 509 496 567 590 625 626 683 671 393 416 451 492 509 496 567 590 625 626 683 671 393 416 451 492 509 496 567 590 625 626 683 671 393 416 451 492 509 496 567 590 625 626 683 671 393 416 451 492 509 496 567 590 625 626 683 671 393 416 451 492 509 496 567 590 625 626 683 671 393 416 451 492 509 496 567 590 625 626 688 629 686 674 393 416 451 492 509 517 307 575 609 630 631 638 676 678 665 407 448 488 503 511 512 569 594 627 628 685 673 396 422 453 386 511 512 569 594 627 628 685 673 396 422 453 386 511 512 512 569 594 627 628 685 673 396 422 453 386 511 512 512 569 594 627 628 686 674 675 694 674 694 694 694 694 694 694 694 694 694 69	369	397	427		4.85			580		495	659	645
372		398			486	347	544	504		520		647
373 323 431 465 489 498 547 587 605 589 603 650 375 336 432 476 490 500 548 588 606 599 664 651 652 652 376 362 434 478 492 505 550 550 554 608 600 666 653 377 363 435 479 493 507 551 555 609 601 667 654 378 364 436 485 494 508 552 556 610 606 668 655 379 366 437 466 495 516 532 556 610 606 668 655 380 368 438 467 496 532 556 554 561 612 621 670 657 381 417 439 469 497 533 555 562 613 369 671 658 382 418 440 470 498 535 556 563 614 393 672 659 383 426 441 473 499 534 555 557 572 615 486 673 660 384 429 442 475 500 468 558 573 616 489 674 661 385 430 443 480 501 348 559 574 617 504 675 662 386 431 444 481 502 399 560 575 618 553 656 633 389 407 447 487 505 442 3562 577 620 591 678 665 389 407 447 487 505 432 562 577 620 591 678 665 399 410 448 488 506 454 564 583 622 597 680 668 391 413 449 490 507 456 565 561 576 619 571 677 664 393 410 448 488 506 454 564 583 622 597 680 668 391 413 449 490 507 456 565 561 576 623 623 623 623 681 669 393 416 451 492 509 496 567 590 625 626 626 627 628 685 673 396 422 453 386 437 445 442 450 591 508 464 564 583 622 597 680 668 391 413 449 490 507 456 565 561 576 619 571 678 665 392 414 450 491 508 464 566 581 624 624 624 682 670 393 416 451 492 509 496 567 590 625 626 627 628 683 671 396 422 453 386 437 513 515 571 602 629 630 688 673 396 422 453 386 437 518 512 509 496 567 590 625 626 627 628 685 673 396 422 453 386 511 512 569 594 627 628 685 673 396 422 453 386 511 512 569 594 627 628 685 673 396 422 453 386 511 512 569 594 627 628 685 673 396 422 453 386 511 512 569 594 627 628 685 673 396 422 453 386 511 512 569 594 627 628 686 674 673 399 436 457 499 515 536 577 600 633 631 632 689 677 640 438 438 438 503 516 537 571 602 629 630 687 675 604 444 440 459 509 517 307 575 601 633 634 691 681 690 444 444 462 519 509 444 578 578 607 633 634 691 681 690 444 444 444 462 519 520 442 578 607 633 634 691 681 690 680 444 444 462 519 520 442 538 577 601 633 634 691 681 400 444 444 462 519 520 544 538 579 619 637 637 639 695 685 685 685 685 685 685 685 685 685 68				439	488	360	546	586	604	564	662	648
375 345 433 477 491 501 549 350 007 590 605 666 653 377 363 434 478 492 505 550 554 608 600 666 653 377 363 435 479 493 507 551 555 609 601 6667 654 668 655 379 366 437 466 495 516 553 560 611 614 669 656 380 368 438 467 496 532 554 561 612 621 670 657 381 417 439 469 497 533 555 566 613 369 671 658 382 418 440 470 498 535 556 563 614 393 672 658 384 429 442 475 500 468 558 573 616 489 674 661 385 430 443 480 501 348 559 574 617 504 675 662 386 431 444 481 502 399 560 575 618 553 676 663 387 448 449 440 470 4498 535 566 575 618 553 388 437 446 484 504 423 562 577 620 591 678 665 389 407 447 487 505 432 563 582 621 595 679 686 393 413 444 450 490 507 456 565 302 623 623 681 669 392 414 450 491 508 464 566 581 624 624 682 670 393 410 448 488 506 454 564 583 622 597 680 668 391 413 449 490 507 456 565 302 623 623 681 669 392 414 450 491 508 464 566 581 624 624 682 670 395 422 453 386 511 508 464 566 581 624 624 682 670 395 422 453 386 511 502 509 496 567 590 625 626 683 671 398 428 456 497 514 531 572 604 630 631 688 676 398 428 456 497 514 531 572 604 630 631 688 676 399 436 457 499 515 536 573 605 631 632 689 686 674 490 438 458 509 517 571 602 629 630 687 673 398 428 456 497 514 531 572 604 630 631 688 676 673 398 428 456 497 514 531 572 604 630 631 688 676 673 398 428 456 497 514 531 572 604 630 631 688 676 673 398 428 456 497 514 531 572 604 630 631 688 676 673 398 428 456 497 514 531 572 604 630 631 688 676 673 398 428 456 497 514 531 572 604 630 631 688 676 673 398 428 456 497 514 531 572 604 630 631 688 676 673 398 428 456 497 514 531 572 604 630 631 688 676 673 398 428 456 497 514 531 572 604 630 631 688 676 673 398 428 456 497 514 531 572 604 630 633 634 691 681 400 438 458 503 516 537 574 607 632 633 634 691 681 400 438 458 509 517 518 518 570 607 633 633 634 691 681 400 438 458 509 517 518 511 570 602 629 630 687 679 682 682 689 677 684 400 438 458 509 517 518 511 570 604 633 633 634 691 681 400 444 444 462 519 509 517 575 619 637 633 633 634 691 681 400 444 444 462 519 509 517 578 617 619 637 639 695		323		465	489	498	547	587	605	589	663	
376 362 434 478 492 505 550 554 608 600 666 653 674 675 663 379 366 437 466 495 516 553 560 611 606 668 655 379 366 437 466 495 516 553 556 610 614 669 656 657 380 368 438 467 496 532 554 561 612 621 670 657 381 417 439 469 497 533 555 556 613 369 671 657 382 418 440 470 498 535 555 562 613 369 671 658 383 426 441 473 499 534 557 572 615 486 673 660 384 429 442 475 500 468 558 573 616 489 674 661 385 430 443 440 501 348 559 574 617 504 675 662 386 437 446 484 504 423 562 577 620 591 678 664 387 433 445 483 503 405 561 576 619 571 677 664 388 437 446 484 504 423 562 577 620 591 678 662 390 410 448 488 506 454 564 583 622 597 680 668 391 413 449 490 507 456 563 583 622 597 680 668 391 413 449 490 507 456 565 581 624 624 682 670 393 410 448 488 506 454 564 583 622 597 680 668 391 413 449 490 507 456 565 581 624 624 682 670 396 422 453 386 511 512 569 594 627 628 683 671 688 673 686 673 396 422 453 386 511 512 569 594 627 628 683 671 688 673 694 640 438 458 503 511 512 569 594 627 628 683 671 688 673 399 436 457 499 515 531 572 604 633 633 634 691 681 400 438 458 503 517 512 569 594 627 628 683 671 684 672 398 428 456 497 514 513 515 571 602 629 630 687 673 398 428 456 497 514 513 515 571 602 629 630 687 673 399 436 457 499 515 536 573 605 631 632 689 667 673 399 436 457 499 515 536 573 605 631 632 688 670 399 436 457 499 515 536 573 605 631 632 688 670 674 440 448 458 503 511 512 569 594 627 628 683 671 688 676 674 440 449 459 509 517 571 572 604 630 631 688 676 674 440 449 459 509 517 571 572 604 630 631 688 676 674 440 440 459 509 517 518 311 572 604 635 633 634 691 681 400 438 458 503 510 511 568 573 605 631 632 689 677 682 682 640 444 440 459 509 517 518 311 576 611 634 635 636 693 683 404 444 444 462 519 520 442 578 578 617 633 635 636 693 683 640 444 444 462 519 520 442 578 578 617 633 635 636 693 683 640 444 444 462 519 520 442 578 579 619 637 639 695 685 685 685 685 685 693 683 694 444 444 445 462 519 520 442 578 579 619 637 639 695 685 685 685 685 685 685 685 685 685 68		336		476				588		599	665	652
377 363 435 479 493 507 551 555 609 601 667 654 655 378 366 437 466 495 516 553 556 610 606 668 655 656 380 368 438 467 496 532 554 561 612 621 670 657 381 417 439 469 497 533 555 562 613 369 671 658 382 418 440 470 498 535 555 562 613 369 671 658 384 429 442 475 500 468 558 573 616 489 674 661 385 430 443 480 501 348 559 574 617 504 675 662 386 431 444 481 502 399 560 575 618 553 676 663 387 446 484 504 423 562 577 620 591 678 665 389 407 447 487 505 432 563 582 621 595 679 667 390 410 448 488 506 454 459 491 508 464 504 429 420 452 493 510 507 456 565 381 624 624 682 670 393 416 451 492 509 496 567 590 625 626 623 623 681 669 395 422 453 386 511 512 568 567 596 622 629 630 687 675 399 436 457 453 494 513 515 571 571 572 628 685 673 399 426 451 492 509 496 567 590 625 626 628 629 686 674 427 395 422 453 386 511 512 568 592 626 626 627 684 675 399 426 452 493 510 511 568 592 626 626 627 684 675 399 426 452 493 510 511 568 592 626 626 627 684 675 399 426 452 493 510 511 568 592 626 626 627 684 675 399 426 457 499 515 533 515 571 602 629 630 687 675 399 436 457 499 515 533 515 571 602 629 630 687 675 399 436 457 499 515 536 573 605 631 632 689 677 399 436 457 499 515 536 573 605 631 632 689 677 399 436 457 499 515 536 573 605 631 632 689 677 399 436 457 499 515 536 573 605 631 632 689 677 399 436 457 499 515 536 573 605 631 632 689 677 399 436 457 499 515 536 573 605 631 632 689 677 399 436 457 499 515 536 573 605 631 632 689 677 399 436 457 499 515 536 573 605 631 632 689 677 399 436 457 499 515 536 573 605 631 632 689 677 399 436 457 499 515 536 577 609 633 634 691 681 400 438 458 503 516 537 575 609 633 634 691 681 400 444 460 517 518 311 576 611 634 635 692 689 677 694 684 400 443 460 517 518 519 520 442 578 617 635 635 693 683 690 680 680 680 680 680 680 680 680 680 68	375	345		477				554		600	666	653
379 366 437 466 495 516 553 560 611 614 669 656 380 368 438 467 496 532 554 561 612 621 670 658 381 417 439 469 497 533 555 562 613 369 671 658 382 418 440 470 498 535 556 562 614 393 672 659 383 426 441 473 499 534 557 572 615 486 673 660 384 429 442 475 500 468 558 573 616 489 674 661 385 430 443 480 501 348 559 574 617 504 675 662 386 431 444 481 502 399 560 575 618 553 676 663 387 433 445 483 503 405 561 576 619 571 677 664 388 437 446 484 504 423 562 577 620 591 678 665 389 407 447 487 505 432 563 582 621 595 679 667 390 410 448 488 506 454 564 564 583 622 597 680 668 392 414 450 491 508 464 566 581 624 624 682 670 393 416 451 492 509 496 567 590 625 626 627 684 672 396 425 453 386 511 512 569 594 627 628 685 673 396 425 453 386 511 512 569 594 626 627 684 672 398 428 456 497 514 531 572 604 633 631 632 689 674 440 440 459 509 515 536 573 605 631 632 689 674 681 403 444 440 459 509 517 514 570 596 628 629 630 687 675 398 428 456 497 514 531 572 604 633 631 632 689 674 440 440 459 509 517 518 511 570 602 629 630 687 675 398 428 456 497 514 531 572 604 633 631 632 689 674 400 438 458 503 516 537 574 607 632 633 634 691 681 403 444 460 517 518 519 575 609 633 634 691 681 403 444 460 517 518 519 575 609 633 634 691 681 403 444 460 517 518 519 520 442 578 611 634 635 692 682 682 682 683 683 690 680 680 684 644 645 645 645 657 559 669 633 634 691 681 684 609 684 600 635 631 632 689 677 684 607 632 633 634 691 681 600 688 676 600 688 676 600 688 676 600 688 676 600 688 676 600 688 676 600 680 680 680 680 680 680 680 680 68	377	303	435	479		507	55 I	555	609	601	667	654
380 368 438 467 496 532 554 561 612 671 670 657 658 381 417 439 469 497 533 555 562 613 369 671 658 383 426 441 473 499 534 557 572 615 486 673 660 384 429 442 475 500 468 558 573 616 489 674 661 385 430 443 480 501 348 559 574 617 504 675 662 386 431 444 481 502 399 560 575 618 553 676 663 387 433 445 483 503 405 561 576 619 571 677 664 389 407 447 487 505		364	436	485	494	508		556			660	655
381	379	366	437	465				561			670	657
382 418 440 470 498 535 556 563 614 393 672 659 383 426 441 473 499 534 557 572 615 486 673 660 384 429 442 475 500 468 558 573 616 489 674 661 385 430 443 480 501 348 559 574 617 504 675 662 386 431 444 481 502 399 560 575 618 553 676 663 387 433 445 484 504 423 562 577 620 591 678 665 389 407 447 487 505 432 563 582 621 595 679 667 389 407 448 488 506 454	381	417		469	490	533	555	562	613	369	671	658
383 426 441 473 499 534 557 572 615 460 973 661 385 429 442 475 500 468 558 573 616 489 674 662 385 430 443 480 501 348 559 574 617 504 675 662 386 431 444 481 502 399 560 575 618 553 676 663 387 433 445 483 503 405 561 576 619 571 677 664 388 437 446 484 504 423 562 577 620 591 678 665 389 407 447 487 505 432 563 582 621 595 679 667 390 410 448 488 506 454 564 583 622 597 680 688 681 <td>382</td> <td>418</td> <td></td> <td></td> <td>498</td> <td>535</td> <td>556</td> <td>563</td> <td>614</td> <td>393</td> <td>672</td> <td>659</td>	382	418			498	535	556	563	614	393	672	659
385 430 443 480 501 348 559 574 617 504 675 663 386 431 444 481 502 339 560 575 618 553 676 663 388 433 445 484 504 423 562 577 620 591 678 665 389 407 447 487 505 432 563 582 621 595 679 667 390 410 448 488 506 454 564 583 622 597 680 668 391 413 449 490 507 456 565 302 623 623 681 689 392 414 450 491 508 464 566 581 624 624 682 670 393 416 451 492 509 496	383	426		473	499	534	557		615	480	674	661
386 431 444 481 502 399 500 575 618 553 677 664 387 433 445 483 503 405 561 576 619 571 677 664 388 437 446 484 504 423 562 577 620 591 678 665 389 407 447 487 505 432 563 582 621 595 679 667 667 390 410 448 488 506 454 564 583 622 597 680 668 391 413 449 490 507 456 565 380 622 597 680 668 392 414 450 491 508 464 566 581 624 682 670 392 424 453 386 511 512	384			475	500	248	550				675	662
387 433 445 483 503 405 561 576 619 571 677 604 388 437 446 484 504 423 562 591 678 665 389 410 447 487 505 432 563 582 621 595 679 666 390 410 448 488 506 454 564 583 622 597 680 668 391 413 449 490 507 456 565 302 623 623 681 669 392 414 450 491 508 464 566 581 624 624 682 670 393 416 451 492 509 496 567 590 625 626 683 671 394 420 452 493 510 511 568 592	386			481		399	560	575			676	663
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726	723	784	767	842	835	900	896	958	1131	1016	971
727	724	785	774	843	848	901	897	959	1153	1017	973
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1055	1012	1081	1035	1107	1060	1133	1071	1159	1187	1185	1146
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Toadstools and Mushrooms and other Larger Fungi of South Australia. By JOHN BURTON CLELAND, M.D. Pp. 178 with 6 coloured plates and 35 text-figures. Adelaide, British Science Guild. 5s.

Those who study Agarics in this country doubtless appreciate the difficulties of workers in distant lands. Many species occur all over the temperate regions of the globe and several of them are readily recognised from illustrations. The additional greater precision of modern descriptions enables some determinations to be made with complete satisfaction. But there are endless difficulties. Some of the earlier collections sent to Europe were named in what would now be regarded as a rather casual manner. As a result many European species were recorded which were really doubtful and on the other hand some species already described from Europe were given new names. Thus an Australian mycologist, for example, has not only to recognise European species but has also to determine what the Australian forms are which appear in the old lists with European names.

Dr Cleland's handbook shows much evidence of these difficulties but he has based his descriptions on the Australian forms and thus it will be possible with further knowledge to determine the identity of the fungus described, no matter what name is now given to it. A large proportion of the 279 species described have been examined by the author and a considerable number are given with his name

as the authority-but there are no Latin diagnoses.

The book opens with a general introduction of 28 pages which deals with points of interest—edibility, luminosity, habitats, uses and the like—which gains from having reference to Australian fungi. This is followed by a glossary. Classification is then outlined and a key is given to the genera. The arrangement of the body of the work follows Rea and indeed the generic descriptions are taken verbatim from British Basidiomycetae for all genera there listed; the additional genera are

based in the main on Killerman (Engler & Prantl).

The handbook will prove of considerable value to mycologists interested in distribution because of the indication it gives of the species of Agarics found in South Australia. It will moreover doubtless do much to stimulate the study of the group amongst those for whom it was primarily written. It forms one of the series of Handbooks of the Flora and Fauna of South Australia whose preparation has been undertaken by the South Australian Branch of the British Science Guild, and the State Government prints the work and publishes it at cost price: the present volume costs less than four shillings in English money. It is to be hoped that such a liberal-minded governmental attitude towards biological science will meet with success in every way and encourage other legislators to act similarly.

NOTES ON CERTAIN CHANGES IN NOMEN-CLATURE IN THE SECOND EDITION OF THE LIST OF COMMON NAMES OF BRITISH PLANT DISEASES

By E. M. WAKEFIELD AND W. C. MOORE

A STANDARDISED "List of common names of British plant diseases" was published in the Society's *Transactions* in 1929 (XIV (1929), 140-77). Recently this list has been revised* and now contains the names of seven additional hosts and some fifty additional diseases.

In this second edition of the list only a few changes have been made in the common names of the diseases, but considerable revision of the scientific names of the relevant pathogens has been found necessary both to bring matters into line with more recent research and in order to ensure conformity to the International Rules of Botanical Nomenclature as revised at Cambridge in 1930. The present article provides explanations for most of the nomenclatorial changes found necessary, and the notes are arranged in groups corresponding to the main groups under which the host plants are arranged in the list. It may be mentioned here, however, that in naming species of Fusarium Wollenweber's (68) classification has been followed throughout. On the other hand, since the classification and nomenclature of bacteria is at present under consideration by the International Society for Microbiology, no attempt has been made to adhere strictly to any one system of bacterial nomenclature, and the names commonly used in this country have been adopted in all instances. Many of the additions denote diseases that were not known to occur in this country at the time the list was originally prepared. Certain other diseases have now been included on account of their increased prevalence during recent years or because the host involved has come more into prominence commercially.

CEREALS

The name *Ustilago Hordei* was used for Covered Smut of barley by Lagerheim (37) in 1889, and since Kellerman and Swingle (35) did not use this combination until the following year, (Pers.) Lagerh. and not (Pers.) Kellerm. & Sw. must be cited as the authority for the name.

Ustilago Tritici presents a similar but rather more difficult case. In

* It has been published as a separate pamphlet and is obtainable from the University Press, Cambridge.

the Second Annual Report for 1889 of the Kansas Experimental Station, appearing in 1890, Kellerman and Swingle ((35), p. 261) first published the name U. Tritici (Pers.) Jens., on the basis of information given in a letter received from Jensen and dated January 24th, 1890. This combination has been quoted by most later authors and was adopted in the first edition of the list. As pointed out by Liro (40), however, the same combination was also made by Rostrup (58) on March 31st of the same year. The Kansas Report for 1889 bears only the date 1890, but in the Report of the Council given in the Kansas Third Annual Report for 1890 (1891), pp. vii-xii, it is stated that "the second annual report (1889)...was issued from the press of the Kansas Publishing House in June". Rostrup's article therefore antedates that of Kellerman and Swingle, and for this reason the name U. Tritici (Pers.)

Rostr. is now cited.

The double citation, (Bri. & Cav.) Eid., used for Helminthosporium Avenae in the first edition was a mistake which appears to have originated in a paper by Ravn(56) and was not detected at the time the list was prepared. This fungus was originally named as var. Avenae-sativae of Helminthosporium teres by Briosi and Cavara (8). When Eidam (23) raised it to the rank of species he used the epithet Avenae, and not Avenae-sativae. Article 58 of the Rules states that when the rank of a group is changed the earliest legitimate name or epithet given to the group in its new rank is valid, provided it is not a later homonym. Accordingly the correct name for the fungus as a species is Helminthosporium Avenae Eid., and this cannot be discarded in favour of H. Avenae-sativae (Bri. & Cav.) as was done by Lindau (1910, in Rab. Krypt.-Fl. ed. 2, Bd. 1, Abt. 9, p. 35). Ravn's use of (Bri. & Cav.) in parenthesis for the epithet Avenae was either a lapsus calami or due to a misunderstanding of the significance of brackets in citations of authors' names. Both Pyrenophora Avenae Ito and Pleospora Avenae Schaffn. & Rathschl. have been described as the perfect stage of Helminthosporium Avenae, and the first named has recently been recorded in Britain by Dennis (18).

Diedicke (19), in 1903, merely surmised the existence of a "Pleospora teres" on barley; he did not see it. Hence his combination is not valid, and Drechsler's (22) Pyrenophora teres is the first legitimate name given to the perfect stage. The correct citation is therefore P. teres Drechsl.

and not P. teres (Died.) Drechsl. as previously given.

A disease of oats occurring in Wales and previously believed to correspond to Halo Blight (Bacterium coronafaciens Ch. Elliott) has now been shown by Davies and Jones (16) to be identical with the nonparasitic Grey-Leaf. There is no evidence that true Halo Blight occurs in Britain, and the entry has therefore been deleted.

PULSE

Further studies of Chocolate Spot of broad beans have made it doubtful whether this disease is caused by Bacillus Lathyri Manns & Taubenh. Riker and Riker(57) suggest that a bacterium closely resembling Pseudomonas seminum Cayley might be involved, whilst unpublished work carried out at Cambridge appears to show that the disease may be due to a fungus. Hence the insertion of a (?) before the name Bacillus Lathyri under broad beans and also under sainfoin, vetches and sweet pea. On the other hand, the evidence that Marsh Spot of peas is not a parasitic disease is now very strong

(9, 36, 52): it is therefore classed as non-parasitic.

Until recently all the damage to pea plants associated with Assochyta in this country has been attributed to A. Pisi. It is now known that the three diseases, caused by three distinct species of Assochyta, and first differentiated in America independently by Linford and Sprague (39) and by L. K. Jones (33), all exist in Britain, although their relative importance has not yet been determined. The true A. Pisi Lib. produces leaf, stem and pod spotting only, and the name suggested for the disease it causes is Leaf and Pod Spot. The other two species (Mycosphaerella pinodes (Berk. & Blox.) Stone* and Assochyta pinodella L. K. Jones) affect the plant in much the same way, but, in addition, they may attack the base of the stem, inducing a Foot Rot. These two species, together with species of Fusarium frequently associated with a stem rot of pea occurring in the west of England (48), are grouped together as the pathogens of a disease to which the name Foot Rot has been applied.

In the first edition Root Rot of peas was ascribed to Aphanomyces euteiches Drechsl. and to Thielavia basicola Zopf. In 1912 Ferraris (24) applied the name Thielaviopsis basicola to what was considered to be the imperfect stage of the last-named fungus, in order to distinguish it from the ascospore stage. He did not, however, imply the existence of two distinct fungi. In 1925 McCormick (43) brought forward evidence, based on cultural work, which strongly suggested that Thielavia basicola and Thielaviopsis basicola are not genetically connected, although often associated with one another. Since it is the conidial stage that is commonly found associated with rotting of pea roots in Britain the name Thielaviopsis basicola (Berk. & Br.) Ferraris has been selected in preference to Thielavia basicola Zopf, and the opportunity has been taken to distinguish Black Root Rot due to this

fungus from Root Rot due to Aphanomyces.

^{*} The name Niessl was left in the list by an oversight. Niessl called the fungus Sphaerella. The change to Mycosphaerella was made by Stone (Ann. mycol. x (1912), 581).

Ротато

The only significant emendation made in the section dealing with potato diseases is that relating to Spraing. Brown spotting or streaking of the flesh of potato tubers, quite distinct from the brown decay due to blight, has been known in Britain for over half a century, but its true nature and origin have always been, and still are, a matter of considerable uncertainty. In England the malady was at first called Canker or Internal Disease, but later the Scottish term Sprain or Spraing came into more general use, both in this country and in Ireland. Horne (31) clearly distinguished two diseases in this category and called them Blotch or Internal Disease, and Sprain or Streak Disease. In Blotch, larger or smaller, brown or chocolate specks, spots or blotches were distributed throughout the flesh of the tubers, whilst in Sprain the markings were usually arc-like or curved in section and often arranged concentrically. Later, Painc (49) introduced the name Internal Rust Spot to include both these sets of symptoms; and they remained grouped together until quite recently, when Grieve (26) segregated them again. In the present list the name Spraing is retained for the disease exhibiting arc-like lesions and the term Internal Rust Spot is applied to the blotch type of lesion. It should be noted, however, that these two types of lesion occasionally occur together in the same tuber. There is at present no evidence that the allied diseases Pseudonetnecrosis and Net Necrosis occur in this country. Spraing has been attributed by different workers to nonparasitic causes (25), to bacteria (10) and to virus agency (26, 55), and the last-mentioned is adopted provisionally in the list. There is some evidence that Internal Rust Spot may be bacterial in nature (10, 26, 49), but the matter has not yet been satisfactorily elucidated.

ROOT GROPS

Opinions differ as to the appropriateness of the names Finger-and-Toe and Club Root for the disease of Crucifers caused by *Plasmodio-phora Brassicae*. In this country usage is about equally divided between them, but in parts of the Empire (15) Club Root appears to be preferred, and largely on account of this the name Finger-and-Toe has now been replaced by Club Root as a common name under turnip,

cabbage, radish, seakale and wallflower.

In the first edition Soft Rot of turnip and swede was ascribed to two distinct organisms, Bacillus carotovorus L. R. Jones and Pseudomonas destructans Potter. The original cultures from which Potter (53) described his organism as a uniflagellate rod were lost, but two later isolations by Potter, which he believed to be identical with his original isolation, were found by Harding and Morse (28) to consist of an organism with peritrichiate flagella which they regarded as

specifically identical with *Bacillus carotovorus*. Since Potter (54) himself appears to have accepted this interpretation the disease is now attributed to *B. carotovorus* only.

White Spot of turnip, previously ascribed to Cercospora Bloxami Berk. & Br., is now attributed to Cercosporella Brassicae (Fautr. &

Roum.) v. Hoehn.*

VEGETABLES

For many years the Downy Mildew of onions and of shallots has been known as *Peronospora Schleideni* Unger, and this was the name included in the first edition of the list. In 1932, however, Cook(13) endeavoured to prove that it should be called *P. destructor* (Berk.) Caspary. Unfortunately Article 57 of the International Rules, concerning the names of fungi having a pleomorphic life cycle, does not mention the Phycomycetes. Mycologists, however, appear to be agreed that the rule should apply to this group also, and it is therefore necessary to discover the name first given to the "perfect" or

oospore stage.

The conidial stage of the onion Downy Mildew was first described by Berkeley (6), in 1841, as Botrytis destructor Berk., but later, in 1860 (7), he listed it as Peronospora destructor Casp. Unger's name P. Schleideni dates from 1847(66), and similarly applies only to the conidial stage. In 1855 Caspary(11) described oospores of various species of *Perono*spora, but there seems to be no evidence that he ever saw those of the onion fungus, and he himself does not appear to have published the combination P. destructor. Cook (13) maintained that because Berkeley used this name for his fungus, and mentioned in the generic description of Peronospora that the genus produces oospores, therefore the name P. destructor was applied to the perfect stage. The argument is false, however, and it cannot be assumed that Berkeley received a private communication from Caspary stating that he had seen oospores. It seems much more probable that Berkeley merely followed Caspary in including this species in the genus Peronospora on general grounds. In point of fact, three years after publication of the name P. destructor by Berkeley, de Bary (17) wrote of the onion fungus "oosporis ignotis".

As far as can be ascertained the oospores of onion Downy Mildew were first mentioned by W. G. Smith (62) in 1884. He gave an illustration of them and stated that they had been found by Vize in decaying patches on onions that had previously borne the conidial stage. Shipley (61) fully described the oospores, and his illustrations of them were drawn by W. G. Smith. In both instances the name used was P. Schleideniana, a combination made by de Bary (17) in 1863. De Bary's name was nevertheless invalid, for he merely applied it instead

of *P. Schleideni* to the conidial stage, and names cannot be changed arbitrarily. Since W. G. Smith used this combination for the oospore stage, however, it is correct to call the fungus *P. Schleideniana*, but the name must be attributed to W. G. Smith and not to de Bary.

The laws of priority necessitated the change from Colletotrichum oligochaetum Cav. to C. lagenarium (Passer.) Ell. & Hals. for the fungus causing Anthracnose of cucumber. This organism was first described in 1867 by Passerini (50) as Fusarium lagenarium, and it was transferred, first to Gloeosporium and later, in 1893, by Halsted (27) to Colletotrichum as C. lagenarium Ell. & Hals. Meanwhile, in 1889, Cavara (12) had described what is now regarded as the same species, as C. oligochaetum and his name is commonly but incorrectly given to the fungus.

The organism causing Bacterial Spot of lettuce is Bacterium marginale N. A. Brown and, adopting a suggestion by Prof. S. G. Paine, this name is no longer given as a synonym of B. pyocyaneum (Gessard) Lehm. & Neum. (45). Paine has pointed out (in litt.) that when first isolated from affected lettuce B. marginale does not exhibit the deep green colour of B. pyocyaneum; moreover, the last-named is regarded by medical workers as a human pathogen and, until further information is forthcoming, it is desirable to avoid possible exaggeration of the importance of this organism in comparative animal and plant

pathology.

The authority for the name Marssonina panattoniana Berl. is now correctly given as (Berl.) Magn. In 1906 Magnus (42) showed that the name Marssonia (often wrongly spelt Marsonia) was given by Karsten to a Phanerogamic genus in 1861, thirteen years before Fischer chanced upon the same name to designate a fungus. Fischer's name was therefore an invalid one and Magnus replaced it by Marssonina, to which he transferred various species, including M. panattoniana Berl. The use of brackets for Berl., the first author of the epithet panattoniana, is, of course, in accordance with Article 49 of the International Rules, which lays down that when a species is either transferred to another genus, or altered in rank, but without changing its specific epithet, the name of the original author must be cited in parenthesis.

Recent investigations (3) have permitted more precise definition to be made of the virus diseases of tomato, and Spotted Wilt, Mosaic, Yellow Mosaic and Streak are now differentiated. In the past the last-named has often been called Stripe, but it is considered desirable to reserve the name Stripe for the disease ascribed to Bacillus Lathyri Manns & Taubenh. The term Streak as now applied embraces four distinct Streak diseases caused by different viruses but indistinguishable by their symptoms alone. Only two of these, viz. Single-virus Streak and Mixed-virus Streak, are known to occur in this country.

The common name for the disease of cabbage due to Phoma

Lingam (Tode) Desm. has been changed from Stem Rot to Canker to meet a criticism made by Cunningham (15).

Fruit

Changes have been made in the name and authorities of the two Brown Rot fungi Sclerotinia fructigena Schroet. and S. cinerea Schroet. wherever they occur in the list. In 1893, Schröter (59), who contributed the section on fungi to Cohn's Kryptogamen-Flora, discussed the two species Monilia fructigena and M. cinerea and named them Sclerotinia fructigena and S. cinerea respectively. Reference to his descriptions, however, shows clearly that he did so only by analogy, for he stated that the perfect stages of these fungi had not been found. His combinations are therefore invalid.

As pointed out by Harrison (29), the first description of the perfect stage of *Monilia fructigena* appears to be one by Aderhold and Ruhland (2) in 1905, and the correct citation for the common fruitrotting fungus of Europe having buff-coloured fructifications is there-

fore Sclerotinia fructigena Aderh. & Ruhl.

In the same article Aderhold and Ruhland described two other fungi in their perfect stages, viz. Sclerotinia laxa and S. cinerea. The account of S. laxa was based upon European material, and this would appear to be the first description of the perfect stage of what has long been known in Europe as Monilia cinerea Bon. The strains of this fungus that cause Brown Rot, Blossom Wilt and allied diseases of pear, plum, cherry, apricot and peach all appear to be identical, and they should all be included under the name Sclerotinia laxa Aderh. & Ruhl. The strain that causes Brown Rot and Blossom Wilt of apple is apparently a distinct biologic form, known hitherto as S. cinerea f. Mali, but now correctly named S. laxa Aderh. & Ruhl. f. Mali (Worm.) Harrison(29).

Aderhold and Ruhland's description of S. cinerea, on the other hand, was based on material received from America, and these authors were mistaken in assuming that they were dealing with the perfect stage of the European Monilia cinerea Bon. In point of fact they had before them a specimen of the common American Brown Rot fungus, now generally called Sclerotinia fructicola (Wint.) Rehm, which is unknown in this country. Indeed, it is very doubtful whether it occurs normally in Europe, although it was isolated once

some years ago in Holland from an apple fruit.

Cooke (14), in 1866, first described the ascigerous stage of the Apple Scab fungus as Sphaerella inaequalis. The combination Venturia inaequalis, usually attributed to Aderhold (1), was actually made by Winter much earlier and was used in Thuemen's Mycotheca Universalis, Nos. 261, 650 and 1544, issued 1875, 1877 and 1880 respectively. The name was used by Winter in a rather wider sense than it is now,

but that does not affect its validity, and the correct citation is *V. inaequalis* (Cooke) Wint. For precision "emend. Aderh." may be added, since the type of Cooke's *Sphaerella inaequalis* was a form on

Pyrus Aria and not on apple.

With regard to the species of Myxosporium causing Surface Canker of apple, since the mistake is so often made it seems desirable to point out that the specific epithet should be corticola and not "corticolum", as was wrongly used by Edgerton. Latin words ending in -cola, as incola, agricola, terricola, are not adjectives, but nouns of common gender, formed from the stem col. There is no corresponding adjectival form. The specific epithet corticola is thus a noun in apposition, meaning an inhabitant of the bark; it cannot be declined like an adjective. The perfect stage of this fungus has not yet been found in this country, but it is known on the Continent under three different names. Madame Arnaud (4) first described it in France in 1923 as Dermatea corticola Arn. In 1930 Jørgensen (34) encountered what is clearly the same organism in Denmark, but he was apparently unaware of Arnaud's work and regarded his fungus as new to science, naming it Neofabraea corticola. More recently Nannfeldt (47) has considered the genus Neofabraea to be synonymous with Pezicula, and has called the apple fungus *Pezicula corticola* (Jørgens.) Nannf. The correct double citation under *Pezicula*, if the synonymy is correct, should be P. corticola (Arn.) Nannf.

Physalospora obtusa (Schw.) Cooke has been substituted for P. Cydoniae Arn. as one of the causal agents of Leaf Spot of apple, in accordance with recent investigations by Stevens (63), and similarly Plectodiscella veneta Burkh. has been replaced by Elsinoe veneta (Burkh.) Jenk. for Anthracnose of raspberry (32), while Gloeosporium ampelophagum (Pass.) Sacc. has been listed as a synonym of Elsinoe ampelina Shear for Anthracnose of grape vine (60). As regards Rust of black currants, Cronartium ribicola, Sydow (64) has submitted legitimate reasons for citing J. C. Fischer instead of Diet. as the authority for

the name.

Loganberry Rust and Branch Die Back of plums have been deleted from the list on account of their apparent rarity in this country. Rough Scab of apples has also been omitted, since in view of recent research by Moore (46) it is very doubtful whether this name can be applied to any specific disease, or whether the fungus to which it was attributed (Coniothecium chomatosporum Corda) can be regarded as actively parasitic. On the other hand, there are now more cogent reasons for regarding the Sooty Blotch fungus on apples in this country as identical with Gloeodes pomigena (Schw.) Colby, and a correction to this effect has been made.

Recent investigations have helped considerably to elucidate the obscurity regarding Die Back in plum trees. Bacteria were pre-

viously thought to be implicated, and Wormald's work (69) has now revealed the existence of a specific disease caused by *Pseudomonas mors-prunorum* Worm. This disease has therefore been removed from the congeries of undetermined diseases included under the term Die-Back and has been named Bacterial Canker.

ORNAMENTALS

The only change made in the common names under this section is the replacement of the name Stalk Break by the more appropriate term Loose Bud for a non-parasitic condition not infrequently ex-

hibited by hyacinths (51).

The discovery and naming of the perfect stage of the gladiolus Dry Rot organism, Sclerotium Gladioli Massey, enables attention to be drawn to a further point in nomenclature often unappreciated by plant pathologists. În fungi with a pleomorphic life cycle, the name to be used is the earliest given to the state which it has been agreed to call the perfect form, provided that this name is otherwise in accordance with the International Rules (Art. 57). It frequently, indeed usually, happens that the author describing the perfect stage uses the same specific epithet as that of the conidial or sterile form previously known. Many writers cite the name of the author of the conidial stage in parenthesis. For instance, when Drayton (21) found that Sclerotium Gladioli Massey produces a Sclerotinia as the perfect state he wrote its name Sclerotinia Gladioli (Massey) Drayton. Since the Sclerotinia was a fungus described de novo, however, and it was not merely a question of changing an existing name which already had a full description, the case is not quite the same as that provided for in Art. 49. If the ascigerous stage had previously been described as Peziza by Massey, the double citation (Massey) Drayt. would have been correct. The two cases are not parallel, and for this reason the brackets have been omitted here and in all instances where it is known that the first author described only the conidial or imperfect

Phytophthora Richardiae Buism., the causal agent of Root Rot of arum (Richardia), is considered by Ashby (5) to be merely a variety of P. cryptogea and it is now listed as P. cryptogea Pethybr. & Laff.

var. Richardiae (Buism.) Ashby.

As shown by Wollenweber ((68), p. 499) the carnation Leaf Rot fungus described by Höstermann and Laubert in 1921 as *Pseudo-discosia Dianthi* is identical with *Excipulina valtellinensis* Trav. (1903). This fungus is, however, a *Heteropatella* and its correct name is *H. valtellinensis* (Trav.) Wr.

A description of the species of *Penicillium* that causes Storage Rot of gladiolus was published from two sources early in 1928, and in both instances the name *P. Gladioli* was used. A preliminary account

of it was published by McCulloch and Thom(44) in No. 1730 of Science, issued on February 24th, 1928, and these authors were cited as the authorities for the name in the first edition. Machacek(41), however, described the same fungus under the same name in the 19th Ann. Rep. Quebec Soc. Prot. Plants for 1926-7, Quebec, 1927, p. 77, and, in 1930, Thom(65) ascribed priority of publication to Machacek. This view was adopted in the second edition, but subsequently it was discovered that according to Drayton (20) the publication containing Machacek's description was not distributed until February 28th, 1928. The correct authority for the name is therefore McCull. & Thom, as given in the first edition.

Although no change has been made in the name *Botrytis Tulipae* for Fire of tulips, enquiries have been received from time to time as to the reason for citing (Lib.) Lind as the authority instead of (Lib.) Hopk. It may be pointed out, therefore, that when Hopkins (30) made the combination in 1921 he overlooked the fact that Lind (38) had already used the same combination eight years previously.

Under iris Soft Rot Bacillus omnivorus van Hall has been listed as a synonym of B. carotovorus L. R. Jones in accordance with modern opinion, and delphinium Mildew, previously listed in error as Erysiphe Cichoracearum DC., has been correctly named E. Polygoni DC.

Recent unpublished work* in this country has demonstrated that Wilt of carnations, as previously understood, is not a single specific disease due to certain species of Fusarium but is a mixture of two or three distinct diseases. One of these—and perhaps the least important economically—has been differentiated as Die-Back (Fusarium culmorum (W. G. Sm.) Sacc.). The others are grouped together for the time being under the common name Wilt and Stem Rot, attributed to species of Verticillium and Fusarium.

In view of Waterman's (67) conclusions that Coniothyrium Fuckelii Sacc. and C. Rosarum Cooke & Harkn. are identical it was decided to omit Graft Disease as a rose disease distinct from Stem Canker. Since the perfect stage of the fungus has been found in this country Stem Canker is now attributed to Leptosphaeria Coniothyrium (Fuck.) Sacc. The so-called Brand Canker, due to Coniothyrium Wernsdorffiae

Laub., has not yet been found in England.

In the first edition Leaf Spot of viola and violet was attributed in part to Ascochyta Violae Sacc. & Speg. From examination of typical specimens during the past few years, however, it would appear that the fungus commonly causing Leaf Spot of violets in this country does not correspond to the original description of A. Violae Sacc. & Speg. It not infrequently produces two-celled spores and is therefore perhaps correctly to be regarded as a species of Ascochyta, but

^{*} For a brief reference to the results of this research see The Fruit Grower, LXXVII (1934), 179.

in general it seems to agree closely with the description of Phyllosticta Violae Desm., and it is provisionally referred to that species. As a cause of Leaf Spot in violas and pansies P. Violae appears to be uncommon in England, and this entry has accordingly been deleted.

MISCELLANEOUS

Thielavia basicola Zopf is renamed Thielaviopsis basicola (Berk.) Ferraris (see p. 99) under Root Rot of tobacco, and a short list of fungi known to attack lawn grasses in this country has been added. Little attention has hitherto been paid to turf diseases, and for the present no useful purpose would be served by attempting to apply common names to them.

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ON CERCOSPORA BLOXAMI BERK. & BR.

By E. W. MASON

Two fungi are passing in current literature as "Cercospora Bloxami Berk. & Br." The first is especially associated with white spot of turnip leaves in this country, but, beyond this fact, has no apparent connection with Berkeley or Broome or Bloxam. It is represented in the Kew Herbarium by the following specimens: G. H. Pethybridge, Devon, 30. x. 24; N. C. Preston, Newport, Salop, x. 24; W. Buddin, Dorset, 12. iii. 26; E. M. Wakefield, Kingthorpe nr. Pickering, Yorks, 18. ix. 30. This is clearly the fungus that was officially recognised by the Plant Pathology Sub-Committee of this Society in 1929 as Cercospora Bloxami Berk. & Br. Consideration of Roumeguère's Fungi sel. exsic. 5679, preserved in the same herbarium, shows that it is also Cylindrosporium Brassicae Fautr. & Roum., which was described in 1891 on "Brassica Napo-brassica Vilm." and which von Hoehnel in 1924 referred to the genus Cercosporella as Cercosporella Brassicae (Fautr. & Roum.) von Hoehn.

The second is especially associated with cabbage leaves (Brassica oleracea and B. sinensis) in the tropics,* and, if I am correctly informed, has not been collected in this country. It is represented in the Kew Herbarium by C. G. Hansford, No. 1296, on cauliflower, Kampala, Uganda, June, 1930, and in the herbarium of the Imperial Mycological Institute by W. F. Steven, San Juan, Trinidad, 11. iv. 34. This is Cercospora brassiciola P. Henn., as represented at Kew by

Baker's Fungi Malayana, No. 17 (det. Sydow.).

There is not the slightest difficulty in distinguishing these two fungi: Cercosporella Brassicae is white to the naked eye, and, under the microscope, the conidia are typically cylindrical and rounded at both ends. Cercospora brassicicola appears to the eye as little blackish tufts; and each conidium, which typically tapers from the base to the apex, has a broad scar right across its base; associated with this, and quite as characteristic, is the presence of brown conidiophores with corresponding broad scars. The only point at issue is to which of these two fungi the name Cercospora Bloxami should be applied.

The species was described in Ann. Mag. nat. Hist. Ser. 5, IX (1882),

183, as follows:

"1979. Cercospora Bloxami, B. & Br. Maculis orbicularibus pallidis; sporis elongato-fusiformibus utrinque acuminatis multiseptatis.—On decaying leaves of turnips, Twycross, Rev. A. Bloxam. Formerly distributed as Septoria Bloxami."

^{*} It is also well known in the United States.

There is a specimen at Kew from Berkeley's herbarium, labelled by Berkeley "Septoria Bloxami B. & Br." On the envelope is written "on decaying leaves of turnips, Twycross. Unknown to me, A.B." [A. Bloxam]. Three spores have also been figured by the same hand at a low magnification; they appear elongate fusiform, multiseptate (up to 14 cross-septa) and acuminate at the ends. Examination of the specimen shows that the spores figured (and to which the diagnosis undoubtedly refers) are those of the fungus Alternaria Brassicae (Berk.) Bolle*; these are there in abundance, but nothing else is to be found.

This then is clearly the type specimen of "Cercospora Bloxami", which was overlooked for so long because Berkeley had not relabelled his "Septoria Bloxami"; and this specific epithet is accordingly not available for either of the two fungi to which it is being currently

applied.

The correct name for the first appears to be Cercosporella Brassicae (Fautr. & Roum.) v. Hoehn. in Ann. mycol. xx (1924), 193, synon. Cylindrosporium Brassicae Fautr. & Roum. in Rev. mycol. xxii (1891), 81. From a consideration of their diagnoses, Hoehnel also suggested that Cercospora (Cercosporella) albomaculans Ell. & Ev. (1894) and Ramularia Rapae Pim (1897) were synonyms, and (quite rightly) that Cercospora Bloxami Berk. & Br. had multiseptate spores, and was therefore distinct. Authentic material of neither Cercosporella albomaculans nor of Ramularia Rapae is preserved at Kew, but von Hoehnel's surmise is very probably correct.

The correct name for the second species appears to be *Cercospora brassicicola* P. Henn., although in this case it does not seem certain that authentic material has been examined in recent times. The type collection is not preserved in Berlin, but, Prof. Dr E. Ulbrich thinks,

may possibly exist in Tokyo, Japan.

^{*} As cited in "The List of Common Names".

THE SPORULATION OF HELMINTHOSPORIUM AVENAE AND ALTERNARIA SOLANI IN ARTIFICIAL CULTURE

By W. A. R. DILLON WESTON, M.A., Ph.D.

(School of Agriculture, Cambridge)

Sometimes there exists a diversity of opinion as to the ease with which a fungus produces spores in artificial culture. Such is the case with the various species of *Helminthosporium*. In view of conflicting statements, Turner and Millard(1) made a detailed study of a culture of *H. Avenae*. A wide range of media was used to embrace varying carbohydrate and nutrient contents, hydrogen-ion concentration and sterilisation methods. No sporulation occurred on any of the cultures excepting on sterilised oat leaves, and then only sparsely. Shortly after this work was published, Dillon Weston(2) stated that abundant sporulation could be promoted by the irradiation of artificial cultures with a Hanovia quartz mercury-vapour home-model alpine sunlamp, alternating current equipment 200 volts.

It is the purpose of this note to indicate the salient points of this work, and to show how sporulation of the fungus causing Early Blight, *Alternaria Solani*, may be induced by similar methods.

HELMINTHOSPORIUM AVENAE

After it had been found that this fungus would sporulate when irradiated with an artificial source of light such as a quartz mercury-vapour lamp, it was necessary to determine the time of irradiation that was necessary at a given distance from the source of light and whether this phenomenon depended upon invisible ultra-violet light

or visible white light.

Non-sporing cultures were made in Petri dishes and also in watch-glasses, with watch-glass tops as covers, and were irradiated for ten minutes at a distance of one foot from the source of light. Some cultures were irradiated through the normal Petri plate covers and others through substituted covers made of a proprietary glass named Sanulux that gave a very considerable transmission at $300 \,\mu\mu$. Sporulation was abundant in these cultures, while non-irradiated cultures made at the same time and under the same conditions failed to spore. This was confirmed by Mr C. C. Brett, of the Official Seed Testing Station, Huntingdon Road, Cambridge. He irradiated non-sporing cultures for five minutes through Sanulux glass with the result that abundant spore formation took place.

The time period was then reduced and cultures were irradiated for periods between one minute and ten, through the ordinary glass cover and through Sanulux glass. At all periods within this range and under both types of glass, sporulation took place. It was noted that the longer periods of irradiation produced an increase of

pigment.

Cultures were also exposed for varying periods to natural irradiation out-of-doors. Some cultures were covered by Sanulux discs, others by the ordinary glass. In all the experiments sporulation took place, but control cultures with blackened surfaces did not sporulate. No attempt was made to determine the time period necessary out-of-doors, since the natural light conditions are variable. On a day during the first week in March, 1933, cultures were exposed for 2 hours 35 min., from 1.55 to 4.30 p.m. Sporulation resulted.

These experiments show that sporulation can be induced by irradiation with natural or artificial light-sources. They suggest, however, that irradiation between 320 and 295 $\mu\mu$ is not the dominant factor in causing spore formation, since it takes place through ordinary

glass.

Experiments were then made to see if spore formation could be brought about if the visible light and the shorter ultra-violet rays were screened off. The lamp was screened with a Hanovia diagnosis filter. This cuts out the visible light and transmits ultra-violet between 400 and 300 $\mu\mu$, the main portion of the ultra-violet being furnished by the line 366 $\mu\mu$. Cultures were irradiated for varying periods up to twenty minutes, but no spore formation took place.

As it seemed that sporulation was brought about by visible light and not ultra-violet light, cultures were irradiated to determine the

shortest period that would be necessary for exposure.

Cultures made in watch-glasses were exposed for varying periods of from one to sixty seconds. In one series they were irradiated through watch-glass covers and in another series with these covers removed. When exposed directly to the artificial light source sporulation was noted at the exposure made for two seconds and at the periods above this. When irradiated through glass covers, spore formation was delayed and was not noted until the exposure had been made for twenty seconds. Cultures were also made in watch-glasses and placed inside a photographic camera, the lens of which had been removed. This was then placed one foot below the artificial light source. Exposures were made by operating the automatic shutter. Different cultures were exposed for 1/25th, 1/50th and 1/100th of a second and also for periods of from one to thirty seconds. No sporulation resulted. As irradiation through a glass cover had increased the time period necessary, and since spores did not develop after cultures were irradiated for thirty seconds through the diaphragm of a camera,

it was inferred that sporulation depended upon the intensity of the

Eight Petri plate cultures were then exposed for fifteen minutes at varying distances from the source of light, the beam being parallel to the cultures. Spore formation took place when the plates were at distances of two, three and four feet-but not further than this. At two feet more spores were formed than at three, and at four feet only a few spores were noted. Controls not irradiated did not spore.

During the course of the experiments the following facts were observed. Spore formation took place about eighteen hours after irradiation. The longer the period that cultures were exposed to light the greater was the degree of pigmentation. If the cultures were not over-exposed the spores germinated normally. If irradiated for long

periods the germination was injured.

ALTERNARIA SOLANI

Potato haulm was received from Dr R. Salaman of the Potato Virus Station, Cambridge, in the summer of 1933. This showed the typical "target-like" lesions of the Early Blight fungus, Alternaria Solani. It was, however, difficult at that date to determine the causal organism because spore formation on the material was sparse. After the material had been kept in a moist chamber for ten days spores were noted, and of these some were seen to belong to species of Alternaria. Cultures were made but few spores developed. Stevens (3) says: "The mycelium grows luxuriantly within the leaf but spores do not usually form until after the death of the supporting tissues when the conidiophores emerge through the stomata or by rupturing the epidermis. Often no spores are formed and rarely are many present."

As it was thought that light might be a factor influencing spore formation, cultures were irradiated. In order to cut out injurious ultra-violet rays a piece of plate glass 1/4 in. thick was placed between the source of light to which the cultures were exposed for ten minutes. Eighteen hours later there was profuse sporulation. Cultures exposed

out-of-doors also sporulated.

DISCUSSION OF RESULTS

It appears that visible white light of a high intensity acts as a stimulus for the formation of spores in Helminthosporium Avenae and Alternaria Solani. If this is so there is some call for the use of an incubator under thermostatic and light control. In the laboratory there must be diminished light intensity, and it may be that this intensity is often not sufficient for the production of spores. An incubator of such a type if built for indoor laboratory use would, of course, require artificial illumination. Alternatively an apparatus could be designed for use out-of-doors in natural light.

SUMMARY

I. Helminthosporium Avenae and Alternaria Solani sporulate abundantly when exposed to a high light intensity.

2. Sporulation is induced by high intensity of visible white light.

3. Continued high light intensities increase the pigmentation.

4. It is suggested that there may be a need for an incubator under thermostatic and light control.

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THE GENUS MILESIA IN GREAT BRITAIN AND IRELAND

By LILLIAN M. HUNTER

Although the type of the genus Milesia was described from Scotland, yet up to the present there are very few records of the species of Milesia for Great Britain and Ireland (Grove, W. B., The British Rust Fungi (1913), Cambridge; Grove, W. B., "The British species of Milesina", J. Bot. LIX (1921), 109-10; Grove, W. B., "Mycological notes. VI", J. Bot. LIX (1921), 311-15; Wakefield, E. M., "The Belfast Foray", Trans. Brit. mycol. Soc. xvII (1932), 5-15; Faull, J. H., "Taxonomy and geographical distribution of the genus Milesia", Contr. Arnold Arb. II (1932), 1-138). Likewise the only published accounts of life histories of species of Milesia from Europe are those of Klebahn (Klebahn, H., "Kulturversuche mit Rostpilzen", Z. PflKrankh. xxvi (1916), 257-77) for M. Blechni (Syd.) Arth. and Mayor (Mayor, "Eug. Notes Mycologiques VIII", Bull. Soc. neuchatel. Sci. nat. Lviii (1933), 23-26) for M. Kriegeriana (Magnus) Arth. from Dryopteris Filix-mas (L.) Schott. It was my privilege during 1933-4 to make certain studies on rusts in England, and one important part of this work was to investigate the life histories of various species of Milesia. A preliminary statement of the results of these investigations on Milesia is included here. Accessory to this work much material was assembled both by personal collecting and through the help of others. These collections add to the species already reported for Great Britain and Ireland and to the stages known there for some of the others.

Inoculation experiments with basidiospores from the teleutospores of *Milesia* species on their fern hosts were made on various *Abies* species (Hunter, L. M., "Preliminary note on Life History Studies of European species of *Milesia*", J. Arnold Arb. XVI (1935)) and spermogonia and aecidia of the following rusts were obtained for the first time:

(1) Milesia Scolopendrii (Fuckel) Arth. (from Scolopendrium vulgare Smith) on Abies alba Mill., and A. concolor Lindl. & Gord.

(2) Milesia Polypodii B. White (from Polypodium vulgare L.) on Abies alba and A. concolor.

(3) Milesia vogesiaca (Syd.) Faull (from Polystichum angulare Presl.) on Abies alba.

(4) Milesia Kriegeriana (Magn.) Arth. [from Dryopteris spinulosa (O. F. Müller) Kuntze] on Abies alba, A. concolor and A. grandis Lindl.

Spermogonia and aecidia were also obtained for Milesia Kriegeriana (from Dryopteris Filix-mas) on Abies alba, and on two new hosts,

A. concolor and A. grandis.

In all of the above-named *Milesia* spp. spermogonia appear from twenty-one to thirty days from the time of inoculation, with an average of twenty-three days. The time from inoculation until the appearance of the aecidia varies considerably for the different species as indicated below:

(1) For M. Scolopendrii: 67-89 days, with an average of 77 days.
(2) For M. Polypodii: 82-89 days, with an average of 86 days.

(3) For M. vogesiaca: 99 days.

(4a) For M. Kriegeriana (from Dryopteris spinulosa): 46-59 days, with an average of 53 days.

(4b) For M. Kriegeriana (from Dryopteris Filix-mas): 60-72 days,

with an average of 64 days.

Accidiospores obtained from culture experiments were used in inoculating various ferns with the result that uredospores were obtained for the following species:

Milesia Scolopendrii on Scolopendrium vulgare.
 Milesia Polypodii on Polypodium vulgare.

(3a) Milesia Kriegeriana (from Dryopteris spinulosa) on Dryopteris Filix-mas, D. spinulosa and D. spinulosa var. intermedia (Muhl.) Underw.

(3b) Milesia Kriegeriana (from Dryopteris Filix-mas) on Dryopteris Filix-mas and D. spinulosa var. dilatata (Hoffm.) Underw.

TT

Hitherto unreported collections of Milesia spp. found in Great

Britain and Ireland are listed below:

Milesia Blechni (Syd.) Arth. on Blechnum Spicant (L.) With.: Kilmun, Argyllshire, Sept. 26, 1932; Benmore Estate, Argyllshire, Sept. 28, 1932 and Innellan, Argyllshire, Sept. 29, 1932, G. D. Darker (II); Holne Chase, Ashburton, Devon, Apr. 19, 1934, L. M. Hunter (II, III); Glendhu, Glencullen, Dublin, Mar. 29, 1934, H. B. S. Montgomery (II, III); Killakee, Dublin, May 19, 1934, P. O'Connor (II, III); Woodfalls Cross, Downton, Hampshire, Nov. 14, 1933, L. Allen and L. M. Hunter (II); Blackdown, Haslemere, Surrey, Dec. 11, 1933, R. Blockey and L. M. Hunter (II).

Milesia carpatica (Wrób.) Faull on Dryopteris Filix-mas (L.) Schott.: Milber Wood, Newton Abbot, Devon, Jan. 1, 1934, L. M. Hunter (II); Milber Wood, Newton Abbot, Devon, Apr. 6, 1934, L. M. Hunter

(II, III).

Milesia Kriegeriana (Magn.) Arth. on Dryopteris Filix-mas. (L.) Schott.: Benmore Estate, Argyllshire, Sept. 28, 1932, G. D. Darker (II); Milber Wood, Newton Abbot, Devon, Jan. 1, 1934, L. M. Hunter (II); Torquay, Devon, Mar. 25, 1934, E. Milton (II, III);

Watcombe Glen, Torquay, Devon, Apr. 8, 1934, L. M. Hunter (II, III); Powerscourt, Wicklow, May 20, 1934, P. O'Connor (II, m).

Milesia Kriegeriana (Magn.) Arth. on Dryopteris spinulosa (O. F. Müller) Kuntze: Benmore Estate, Argyllshire, Sept. 28, 1932, G. D. Darker (II); Duloe, Liskeard, Cornwall, Mar. 31, 1934, M. P. Hall (II, III); Watcombe Glen, Torquay, Devon, Apr. 8, 1934, L. M. Hunter (II, III); Powerscourt, Wicklow, May 6 and 20, 1934, P.

O'Connor (II, III).

Milesia Kriegeriana (Magn.) Arth. on Dryopteris spinulosa var. dilatata (Hoffm.) Underw.: Milber Wood, Newton Abbot, Devon, Jan. 1, 1934, L. M. Hunter (II); Milber Wood, Newton Abbot, Devon, Apr. 6, 1934, and Holne Chase, Ashburton, Devon, Apr. 19, 1934, L. M. Hunter (II, III); Killakee, Dublin, May 19, 1934, P. O'Connor (II, III); Woodfalls Cross, Downton, Hampshire, Nov. 15, 1934, L. M. Hunter (II).

Milesia murariae (Magnus) Faull on Asplenium Ruta-muraria L.: Clonsilla, Dublin, Apr. 24 and May 1, 1934, P. O'Connor (11);

Powerscourt, Wicklow, May 6, 1934, P. O'Connor (11).

Milesia Polypodii B. White on Polypodium vulgare L.: Innellan and Sandbank, Argyllshire, Sept. 29, 1932, G. D. Darker (II); Marldon, Torquay, Devon, Dec. 31, 1933, L. M. Hunter (11); Berry Pomeroy, Torquay, Devon, Apr. 2, 1934 and Marldon, Torquay, Devon, Apr. 23, 1934, L. M. Hunter (II, III); Annahilt, Hillsboro, Down, Aug. 25, 1934, L. M. Hunter (II); Glencullen, Dublin, Apr. 1, 1934, H. B. S. Montgomery (II); Powerscourt, Wicklow, May 6, 1934, P. O'Connor (II, III); Woodfalls Cross, Downton, Hampshire, Nov. 15, 1933, L. Allen and L. M. Hunter (II); Blackdown, Haslemere, Surrey, Dec. 11, 1933, L. M. Hunter (II).

Milesia Scolopendrii (Fuckel) Arth. on Scolopendrium vulgare Smith: Duloe, Liskeard, Cornwall, Mar. 31, 1934, M. P. Hall (II, III); Cockington, Torquay, Devon, Dec. 28, 1933, L. M. Hunter (11); Torquay, Devon, Mar. 25, 1934, E. Milton (11, 111); Bishop's Walk, Torquay, Devon, Apr. 1, 1934, and Watcombe Glen, Torquay, Devon, Apr. 22, 1934, L. M. Hunter (II, III); Glencullen, Dublin, Apr. 1, 1934, H. B. S. Montgomery (II); Hale, Downton, Hampshire, Nov. 14, 1933, L. Allen and L. M. Hunter (II); Powerscourt, Wicklow,

May 20, 1934, P. O'Connor (II, III).

Milesia vogesiaca (Syd.) Faull on Polystichum angulare Presl.: Killakee,

Dublin, May 19, 1934, P. O'Connor (II, III).

Milesia Whitei Faull on Polystichum angulare Presl.: Watcombe Glen. Torquay, Devon, Apr. 8, 1934, and Lincombe Hill, Torquay, Devon, Apr. 23, 1934, L. M. Hunter (II, III); Powerscourt, Wicklow, May 20, 1934, P. O'Connor (II, III).

The uredospore and the teleutospore stages as indicated, of the

The Genus Milesia in Great Britain and Ireland

following species of *Milesia*, are reported for the first time from England and Ireland:

England: M. Blechni (III); M. carpatica (II, III); M. Kriegeriana on

Dryopteris Filix-mas (III); Milesia Polypodii (III); M. Whitei (III).

Ireland: M. Blechni (III); M. Kriegeriana on Dryopteris Filix-mas (II, III); M. Kriegeriana on Dryopteris spinulosa (II, III), Milesia murariae (II); M. Polypodii (III); M. Scolopendrii (III); M. Whitei (II, III).

SOME NEW BRITISH RECORDS OF FUNGI ON WHEAT

CERCOSPORELLA HERPOTRICHOIDES FRON., GIBELLINA CEREALIS PASS., AND OPHIOBOLUS HERPOTRICHUS (FR.) SACC.

BY MARY D. GLYNNE

(Department of Plant Pathology, Rothamsted Experimental Station, Harpenden, Herts.)

1. Cercosporella herpotrichoides Fron., considered one of the most important of the fungi causing foot rot of wheat in certain parts of France (3, 4) and of the United States (10), and recently recorded in Germany (8), Holland (6), and Denmark (5), has this year been found at Rothamsted.

Pale lesions with dark borders were noticed occasionally on leaf blades and more often on the outer sheaths and leaf bases of wheat on Broadbalk field and an adjacent plot in February. By the latter half of March many plants still had only the outer leaves and sheaths affected, but the fungus was also found on dead leaves, dead tillers and even dead plants and appeared to be responsible for the damage. Dark-bordered lesions were observed at the bases of sheaths and culms in April and May, and were obvious at harvest in August. The disease was fairly common and was found on every plot on Broadbalk field where wheat has been grown every year since 1843. It was found occasionally but not commonly on some of the other fields in which wheat is grown in rotation with other crops. This accords with the general view that the disease increases when inadequate rotation is practised.

Material consisting of dark-bordered leaf and sheath lesions, of dying tillers and dying plants collected several times in the latter half of March all produced abundant spores in damp chambers in a few days. Hardly any spores were obtained from similar material collected at intervals subsequently. This suggests that the duration of sporing may be very short or only possible under particularly limited

external conditions.

The spores are hyaline, slightly curved, wider at one end than the other, and attached singly or less often in pairs by their larger ends. They are two- to several-septate, generally five to seven. Measurements of twenty spores showed a variation of $31-80 \times 2\cdot 2-3\cdot 3$ μ with an average of $59 \times 2\cdot 3$ μ . These fall within the limits given by Sprague and Fellows (10) of $30-80 \times 1\cdot 5-3\cdot 5$ μ , most spores being 40-60 μ long.

Cultures from spore suspensions grew on potato dextrose agar at 25° C. first as hemispherical mounds of grey velvety mycelium with a pale edge, later growing out over the agar rather slowly and becoming darker on the under surface. They agreed in appearance with cultures of *Cercosporella herpotrichoides* Fron., obtained from the Centraalbureau voor Schimmelcultures, Baarn, representing isola-

tions by Foëx in France and Oort in Holland.

2. Gibellina cerealis Pass., which causes the "white straw disease" in wheat, was found in May on Hoos field, Rothamsted, in the plot on which wheat has been grown alternately with fallow without manure since 1856. The crop was very thin and poor and had suffered much from attack by wheat bulb fly. The fungus caused rotting of tillers and stunting of shoots. It is characterised by dark-bordered elongated lesions on the lower leaf sheaths and basal parts of the culms with a greyish white mycelial felt penetrating and uniting the leaf sheaths, and developing into a stroma with darker cells below. Numerous pale perithecia with black protruding beaks were embedded in the stroma. Only one or two were ripe at the end of May, but from June onwards ripe perithecia were common on affected plants.

Measurements (ten perithecia) showed a variation of 315–600 \times 285–570 μ with an average of 432 \times 395 μ for the perithecium and 285–455 \times 125–220 μ with an average of 367 \times 159 μ for the beak. There were numerous filamentous paraphyses among the asci which measured (twenty asci) 90–125 \times 13–18 μ with an average of 105 \times 16 μ . The uniseptate spores arranged in two rows in the ascus were hyaline at first, becoming honey coloured to hazel and rarely 2–3 celled. Twenty spore measurements showed a variation of 23–36 \times 7–11 μ with an average of 30 \times 9 μ . Except in the width of the asci these measurements more than cover the range given for Gibellina cerealis Pass. Comparison with No. 3669 Rabenhorst-Winter, Fungi europaei, collected by Passerini (Herbarium, Royal Botanic Gardens, Kew)

indicate that it is the same fungus.

Germination of spores was not observed, and Ferraris (2) states that they take a long time to mature. Cultures made from the mycelium, however, grew on potato dextrose agar at first as white mounds which later sometimes turned pale grey. Numerous perithecia developed both in the medium and in the aerial mycelium producing asci with typical honey coloured spores in cultures about five weeks old.

The disease was recorded by Passerini(7) in 1886 in Northern Italy, and has recently been noted by Ferraris(2) as occasionally appearing in May doing slight damage. As crop rotation is unfavourable to the disease and is practised in Italy he considers the disease unlikely to become prevalent. Sprague(9) reports it on winter wheat and oats in Oregon, where it occurs locally on some red

sandstone-shale soils, acid in reaction, in the humid coastal regions. He found only immature perithecia and supposes that the dry

summers of Oregon are not favourable for their maturation.

3. Ophiobolus herpotrichus (Fr.) Sacc. Ripe perithecia were found in March on wheat stubble that had over-wintered in the soil, but no evidence of parasitism was obtained. The perithecia are black with a conical curved beak; the ascospores needle-shaped, multi-septate, measuring (twenty spores) 123-190 × 2·2 μ with an average of 155 × 2·2 μ. Cultures on potato dextrose agar are whitish grey or brownish, often with dark and light areas. As they become older the under surface becomes darker and ultimately black. The fungus occurs in several European countries together with other fungi causing foot rot of wheat and is generally regarded as a weak parasite of secondary importance. In America(1) it has been found on Agropyron repens (L.) Beauv. but not on cereals. In Great Britain it has also been recorded on wild grasses, but not previously on cereals.

I wish to record my thanks to Miss E. M. Wakefield for help in the identification of these fungi.

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CYLINDROSPORIUM CONCENTRICUM GREV.

By J. R. THOMSON, B.Sc.

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(With 8 Text-figures)

Introduction

In 1823 Greville described a fungus producing white spots on cabbage leaves. It was first discovered in a garden at Parson's Green, near Edinburgh, but Greville remarks: "It does not seem to be very rare as I have noticed it in various localities since and probably few gardens will be found without it at some time or other of the year." Unfortunately he gives no more precise information as to the distribution of the fungus at that time.

For the reception of this fungus Greville founded the genus

Cylindrosporium, with the following characters:

Very minute plants, parasitic on living leaves, without rupturing the epidermis. Sporidia cylindrical, truncate, jointless, naked, free, forming little heaps.

The species he called C. concentricum and described it as follows:

Very white and excessively minute but sufficiently conspicuous from their number and from being disposed in concentric lines. Form most irregular; frequently oblong, lying in the direction from the centre to the circumference, projecting into angles and little eminences and having often one extremity turned up like the end of a canoe. Sporidia very numerous, cylindrical, truncate at both extremities, pellucid without joints and not furnished with any membrane or covering whatever. Turning in decay to a dirty yellow, which always commences in the centre, while new individuals are forming at the circumference. The peculiarity of its cylindrical sporidia, and its situation on the surface of living leaves, fully entitle it to generic distinction.

In 1849 the genus Gloeosporium was founded by Montagne. Into it he put the species previously included in Myxosporium and Asteroma

and several new species.

During this time there seems to have been some doubt about Greville's fungus, and there is no record of anyone except Greville having collected it. In 1850, however, it appeared in abundance in market gardens in Northamptonshire. Berkeley(1) immediately recognised it as Greville's fungus and took the opportunity of examining fresh material. He described the fungus as follows:

The parasite forms, both upon the upper and under surface of the leaf, roundish often confluent patches, varying greatly in size, consisting of little white specks disposed more or less concentrically, those of the centre frequently becoming yellow, and at length fading away, in consequence of the partial decomposition of the leaf which they have affected, while the outer pustules spread from the circumference to the part yet remaining healthy. Occasionally they extend to

the midrib, which is then rapidly destroyed. On close examination it is found that the fungus, each speck forming a distinct individual, is produced between the true cuticle and the cuticular cells....The cuticular cells, however, are much confused and deranged by the growth of the parasite, which is developed principally at their expense, those of the succeeding layer being very little if at all affected. The mycelium is closely incorporated with the cuticular cells, and appears simply grumous, without distinct structure;...it does not appear to be filamentous. From the top of this mass, on the level of the tips of the cells on which it grows, arise very short delicate sporophores, each of which is surmounted by an oblong, cylindric, often curved, spore, three to five times as long as broad, and containing at maturity from two to three globose nuclei. It is highly probable that each sporophore produces in succession several spores, which are thus pushed forward and in time fill the space between the true cuticle and the cuticular cells, thrusting the former out until it bursts. Partly owing to the successive development of the spores, which are mixed with a viscid fluid, and partly to the contraction of the leaf itself upon the pulpy mass, in dry weather the spores ooze out, kept in connection with each other by their attendant mucilage, and drying as they are exposed to the air, form rude irregular short tendrils. These tendrils are in their turn softened again by moisture, and after a time fall down, forming a little pellicle on the leaf, the edges of which are often turned up like a little boat or canoe, as observed originally by Dr Greville. The spores, it should be observed, are not truly truncate, as they appeared to Greville when examined by the old imperfect compound microscope, but rounded and obtuse.

Berkeley considered that Greville was correct in founding a new genus, but that Cylindrosporium was identical with Gloeosporium. He added:

It is but proper courtesy to adopt his (Montagne's) name unless he should think fit to restore that of Greville as to the identity of which there is now no doubt.

It was therefore proposed that the fungus should be called Gloeo-

sporium concentricum Berk. & Br.

Saccardo (6) adopted Berkeley's suggestion, though expressing some doubt about it, and in his *Sylloge Fungorum* Greville's fungus is given as *Gloeosporium concentricum* (Grev.) Berk. & Br. (= Cylindrosporium concentricum Grev.). He said that: "From Greville's figure it appears

to be a doubtful species."

Considerable confusion has arisen as to the genus Cylindrosporium and it is difficult to trace its history. The first to use Greville's name was Unger, who added parasites on the celandine and on Brassica spp. under the impression that they were of the same genus as Greville's fungus. These fungi had filiform spores and it is difficult to understand how this confusion arose, for, though Unger probably never saw Greville's actual specimen, he must have seen his description. When Berkeley removed C. concentricum Grev. to the genus Gloeosporium he realised that Unger's genus was quite distinct from Greville's.

Saccardo (6) gave a clear definition of Cylindrosporium Ung. which might have been expected to prevent any confusion. He defined it as a Gloeosporium with filiform conidia and certainly a conidial stage

of *Entyloma*. Unfortunately, according to Diedicke(3) and von Hoehnel(5), some of the species included by Saccardo were not conidial stages of *Entyloma*, and other species, not *Entyloma* forms, have been added since.

In 1924 von Hoehnel (5) investigated the thirty-three species then attributed to *Cylindrosporium* Ung. em. Sacc. (non Grev.). He found that they could be divided into two groups—Melanconieae species and Sphaeroidiae species, the former including conidial forms of *Entyloma* and *Doassansia*. The species were all removed to other genera, five of which were new.

Having demolished the genus Cylindrosporium Ung., von Hoehnel proposed to re-establish the genus Cylindrosporium Grev. with C. concentricum as its only species. While this attitude may be tenable, the reasons which von Hoehnel gives for adopting it seem rather flimsy,

especially as he had never seen a specimen of the fungus.

Von Hoehnel says: "Dieselben (Berk. & Br.) fanden, dass der Pilz, entgegen der Angabe Grevilles, sich unter der Kutikula wie ein Gloeosporium entwickelt und stellten daher den Pilz in diese Gattung.

Unter Kutikula ist hier wohl die Epidermis zu verstehen."

This refers to a short note (2) on Greville's type specimen, and it is obvious that he cannot have seen Berkeley's paper (1) published a year later (quoted above). His description of the fungus differs from Greville's in only one point—the spores are described as having rounded ends while Greville described them as truncate. Moreover, Berkeley was particularly emphatic as to the position of the acervulus—"between the true cuticle and the cuticular cells"—so that the question of confusion between cuticle and epidermis does not arise.

Ignoring Berkeley and Broome's description, von Hoehnel's conclusion is based on that of Greville which is accepted as being correct. From the habit of the fungus, its snow white fructifications and the probable formation of the spores in loose chains, he concludes that it could not be a *Gloeosporium*. The catenulate arrangement of the spores is deduced from Greville's statement that the spores are truncate, which was denied by Berkeley. It must be admitted, however, that from Greville's inadequate description alone it would be difficult to state definitely that it was not a *Gloeosporium*.

Further, von Hoehnel says: "Greville nennt den Pilz eine sehr ungewöhnliche Pflanze, was er gewiss nicht gesagt hätte, wenn es sich um ein Gloeosporidium gehandelt hätte." But the genera Gloeosporium and Gloeosporidium were not established till many years after Greville's time and it may be presumed that fungi of that type

were not then well known.

INVESTIGATION

A fungus forming white spots on cabbage leaves is fairly common in gardens in the Edinburgh district and can frequently be seen on the outer leaves of cabbages exposed for sale in the town. It is generally assumed to be Gloeosporium concentricum (Grev.) Berk. & Br. Inquiries have shown that its distribution is fairly general in Scotland and England, though nowhere abundant. Cauliflowers and broccoli are also attacked. The type specimen of Greville's Cylindrosporium concentricum is in the herbarium of the Royal Botanic Garden, Edinburgh. It was decided to investigate the fresh material and the type specimen and to find what relation, if any, there was between them.

Fresh material

Fresh material was easily obtained in October. The fungus forms white spots on either surface of the leaf, but more commonly on the under surface. The spots are grouped in concentric circles and each group is from 1 to 2 cm. in diameter (Fig. 1). They appear generally

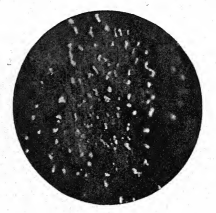


Fig. 1. Group of fructifications on cabbage leaf. $\times 5$.

between the main veins, but may also be found on the veins. As new spots are formed on the outside of the group, the spots in the centre disappear and the leaf turns yellow and then black, and finally only a number of concentric black lines is left. Each group of spots is quite distinct, but the groups may run into each other and overlap. The fungus is found on the outer leaves, which are already beginning to yellow, and seems to have no serious effect on the host. A cabbage which had its outer leaves infected was kept under observation. Some of the adjacent leaves became infected, turned yellow and dropped off, but infection never reached the heart. Finally the fungus dis-

appeared altogether and the cabbage appeared perfectly healthy and very vigorous. The fungus can live saprophytically on dead leaves. As cabbages grow at all times of the year there is no difficulty in the

fungus finding a host.

The spots are minute, being less than 1 mm. across. Under the lens, a spot appears as an irregular gelatinous white mass oozing from a slit in the surface of the leaf. This exudation sometimes takes the form of a tendril, but more generally the spots are irregular and may take the form of a canoe, as described by Greville (Fig. 2).

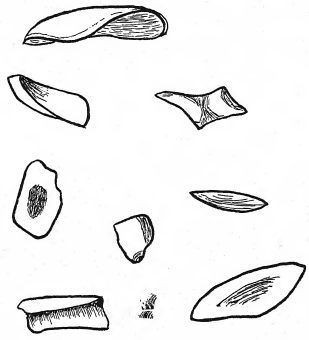


Fig. 2. Fructifications (spots) as seen on leaf surface. × 40.

While the white spots appear to ooze out of a slit in the leaf surface,

there is no apparent rupture or tearing of the epidermis.

Each spot consists of a mass of spores. The spores are cylindrical, sometimes curved, with rounded ends and generally two oil drops. They are $8.5-15\times2.5-5.5\mu$, with an average size of $11\times4\mu$. In January spores were found with an irregular number of transverse bands. This was due to enlargement of the oil drops and the formation of the protoplasm into bands between them (Fig. 3).

Sections were cut of infected leaves. In section the fructification appeared as an irregular grumous mass surmounted by a heap of

spores (Fig. 4). The position of the spores in irregular chains suggested that they were budded off in succession from the basal hyphae, though no section could be obtained showing the process of budding. Where the lamina was infected the tissues were somewhat disorganised, but

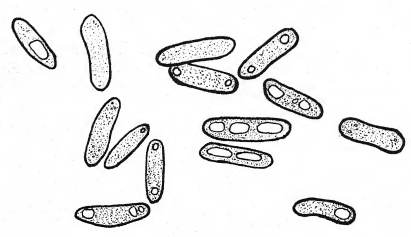


Fig. 3. Spores. × 3600.

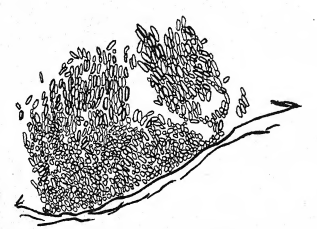


Fig. 4. Section of fructification from leaf lamina. ×360.

the fungus was obviously superficial. Sections through fructifications over a vein, however, showed that they were subcuticular, and no trace of penetration through the epidermal walls could be found (Fig. 5). Sections were obtained showing hyphae within the cuticle between fructifications. The hyphae appeared singly or in pairs, but

occasionally several were clumped together and may have repre-

sented a young fructification (Fig. 6).

It will be seen that this fungus agrees exactly with Greville's description (as far as it goes) of Cylindrosporium concentricum, except for the form of the spores which Greville gave as truncate. This difference, however, was plausibly explained away by Berkeley (see p. 124). It also agrees with Berkeley's description, though no definite

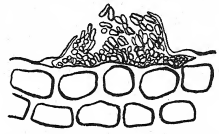


Fig. 5. Section of fructification from vein, showing subcuticular position. × 360.

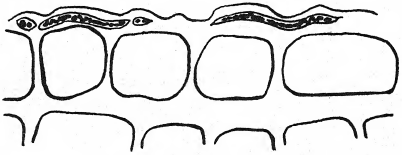


Fig. 6. Section of epidermal cells, showing hyphae within cuticle. × 1270.

sporophores could be found. Berkeley figured the sporophores but does not show them in his figure of the acervulus. This figure shows a flat layer of fungal tissue surmounted by a single layer of spores with no definite attachment. A very similar appearance is seen in hand sections of the fresh material in which the upper spores are knocked off leaving only a single layer.

Saccardo described Gloeosporium concentricum as having truncate spores, and this is the only point of difference between his description

and the fungus examined.

There can be no doubt that this fungus and Greville's are identical.

Type specimen

The type specimen bears the legend "Cylindrosporium concentricum nov. gen. mihi. On Brassica oleracea. Balmuto [Fife], May 1822. Miss Boswell". It consists of a small piece of cabbage leaf with pale-coloured areas but showing no external signs of the fungus.

Sections were cut and fructifications were found. They were quite similar to the fructifications on fresh material, but flattened out owing to herbarium treatment (Fig. 7). Owing to the disorganisation of



Fig. 7. Section of fructification from Greville's type specimen. × 360.

the tissues it was impossible to determine their exact position, but they appeared to be superficial. Free spores could not be obtained so the spores were measured in the sections and compared with spores in sections of the fresh material after similar treatment:

Type specimen Fresh material Spore size range $5 \cdot 5 - 8 \cdot 0 \times 1 \cdot 6 - 2 \cdot 5 \mu$ $5 \cdot 5 - 8 \cdot 0 \times 1 \cdot 6 - 2 \cdot 2 \mu$ Average $6 \cdot 6 \times 2 \mu$ $6 \cdot 6 \times 2 \mu$

It should be noted that the spores are not truncate but have rounded ends. There is a co-type specimen in the Kew Herbarium and free spores have been obtained from it. These spores show this character very clearly.

This investigation of the type specimen confirms the conclusion that the fungus found on the cabbage and Greville's fungus are

identical.

Cultures

Spores from fresh material were germinated in water and dilute cabbage extract. They germinated by a single terminal or sub-

terminal germ tube.

Attempts to culture the fungus on malt and oatmeal agar failed. The spores germinated but soon died off. A medium was therefore made up with cabbage extract. 200 gm. of fresh cabbage leaves were boiled in water for $7\frac{1}{2}$ hours, the extract filtered and made up to 200 c.c. The best results were obtained on a medium containing 5 per cent. of this extract.

The fungus grows very slowly, ultimately forming a growth 1-2 cm.

in diameter. The mycelium, which is greyish, grows within the medium, but frequently a white cottony mycelium is produced on the surface. Later, white gelatinous pustules appear, very like the spots on the cabbage leaf. These pustules consist of heaps of spores as on the cabbage.

The spores are budded off irregularly from the hyphae, there being no specialised sporophores. They are produced laterally or terminally

in succession and lie loosely in irregular chains (Fig. 8).

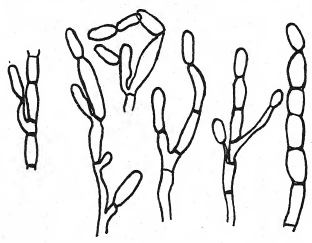


Fig. 8. Spore formation in culture. × 1800.

At a late stage small black sclerotium-like bodies may be produced in culture. They are spherical or lobed and no definite internal structure could be made out. They may possibly be immature pycnidia, but no spores could be found in them. In its final form, black radiating lines appear in the growth giving it a dark appearance, but the periphery remains grey.

DISCUSSION

The question arises as to the systematic position of the fungus. Berkeley and Broome regarded it as a Gloeosporium and were followed in this by Saccardo. The alternative is to retain Greville's genus Cylindrosporium and to regard it as monotypic as suggested by von Hoehnel. Certainly there is nothing in the description of the genus Gloeosporium as given by Saccardo which definitely excludes the fungus from that genus, except the subcuticular position of the accervulus, and Saccardo included it in spite of this. On the other hand, it is certainly not a typical Gloeosporium. The heaping up of the spores

in clumps is unusual, and there seems to be no record of another subcuticular species in the genus. Gloeosporium species have typically a shallow fructification bearing a shallow layer of spores, while in this fungus the spores appear to be budded off in chains and adhere together in a heap above the fructification. Cultures have shown that this arrangement of the spores is due to their being budded off in succession from the hyphae. von Hoehnel thought that its habit, the whiteness of the fructifications, and the probable arrangement of the spores in chains (deduced from the truncate spores) were sufficient to distinguish this fungus from Gloeosporium. While these characters alone seem inadequate, when they are combined with the subcuticular position of the acervulus, the formation of the spores by irregular budding, and the confirmation of the catenulate arrangement of the spores (though they are not truncate), there seems good reason for retaining Greville's genus Cylindrosporium.

Summary

Cylindrosporium concentricum Grev. was described by Greville in 1823. Berkeley removed it to the genus Gloeosporium Desm. & Mont., and was followed in this by Saccardo.

von Hoehnel in 1924 removed all the species from Cylindrosporium Ung. em. Sacc. (non Grev.) and re-established the genus Cylindro-

sporium Grev. though he had not seen Greville's fungus.

A fungus parasitic on cabbage leaves is described and compared with the type specimen of Greville's fungus. It is shown to be identical with the type specimen and to agree in external characters with Greville's description.

Cultures of the fungus were obtained.

The fungus is not a typical Gloeosporium. The description appears to support von Hoehnel's view that the genus Cylindrosporium Grev. should be re-established with this fungus as its type species.

I am indebted to Dr Malcolm Wilson for suggesting this investigation and for his supervision.

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STUDIES IN THE GENUS USTULINA WITH SPECIAL REFERENCE TO PARASITISM

II. A DISEASE OF THE COMMON LIME (TILIA VULGARIS HAYNE) CAUSED BY USTULINA

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(With Plate I and 8 Text-figures)

I. Introduction

As stated in a previous communication (26), the main objective of these studies is investigation into the economic aspect of the temperate form of *Ustulina* in relation to standing timber—especially beech. Beech was chosen because the timber is important and because, in my opinion, *Ustulina*—which is characteristically abundant on that tree—causes a timber rot which is sufficiently serious to merit attention.

Under the general heading, however, the above-mentioned disease of lime is a specific case which occurred locally, and, as it affords excellent material for relevant investigation as well as opportunity for testing experimental methods, will be considered first and is the only aspect of the problem dealt with in the present paper.

Lime is characteristically immune to any form of fungus disease and particularly so to disease of the "timber rot" type; it is not therefore surprising to find that there appear to be only six previous records of *Ustulina* on this host. The first three of these, Lind (13) from Denmark, Bizzozero (1) from Italy, and Sydow (22) from Germany, merely indicate growth of the fungus on the tree with no suggestion of parasitism. Of the last three, however, Wehmer (24), in Germany, is doubtful whether or not it causes disease; Van Overeem (23) says that it occurs in France, is emphatic that it causes disease of lime in that country, but gives no circumstantial details; while Patouillard (16), from France, definitely states that *Ustulina* is parasitic. He says*:

Ce champignon bien connu à l'état saprophyte paraît avoir causé la mort de deux troncs de Tilleul dans l'Ain, en attaquant par la base, au niveau du sol. Sur une hauteur de 20 à 30 cent. ces deux troncs étaient recouvert d'une couche continue du parasite, sauf sur une largeur de 10 cent. qui est restée saine.

Le bois est complètement envahi par le mycelium et a pris une consistance très

* This important reference does not appear in the literature list given in Part I, as it was verified too late to be included in that paper.

molle, jusqu'au centre des arbres, en sorte qu'une simple coup de vent a pu les faire tomber.

L'Ustulina est très fréquent sur les vieilles souches voisines tant sous sa forme ascophore qu'à l'état conidifère mais toujours en saprophyte.

Dans le cas actuel il est nettement parasite.

This scarcity of previous record of disease of lime by this fungus does not entirely eliminate the possibility of its being a pathogen of economic importance as the disease may have been overlooked—a factor which I should like to suggest is also largely responsible for its non-recognition as a pathogen on beech, though this is even less understandable. A second and more probable reason is that the disease symptoms of *Ustulina* may have been credited to another fungus of the "black-line" type.

No standard text-book on the diseases of trees from the time of Hartig (8) in 1882, to the Forest Pathology of Hubert (12) published in

1931, makes any reference to Ustulina as a wood destroyer.

I have come across three lime trees which appear to have been diseased by *Ustulina*:

(1) A tree brought down by the wind in the Botanic Gardens at Oxford.

(2) The tree next to the above which is obviously infected but is still standing.

(3) A very diseased tree in Blenheim Park, Woodstock. This tree is quite hollow, shows all the signs of *Ustulina* disease, and no fructifications other than those of this fungus were found on the tree.

Of these trees only the first mentioned was available for experi-

mental purposes.

II. PRELIMINARY EXAMINATION OF THE TREE

The tree was a well-grown specimen about seventy years old, and, apart from a few dead branches, the crown appeared to be perfectly healthy. Closer examination revealed the fact that the trunk was less sound, there was a wound on the north side—caused when the tree was struck by lightning about 1880—extending from ground level to a height of about twelve feet. The exposed wood of the wound was soft and rotten, but the rest of the trunk seemed to be perfectly normal. There were no sporophores or visible fructifications of any fungus except those of the saprophytic *Peniophora quercina* (Pers.) Cke. on some of the dead branches, and those of *Ustulina vulgaris* Tul. which were growing on the surface of the exposed wood of the wound. There was no evidence of the rhizomorphs of *Armillaria mellea*.

When the tree crashed the base was badly smashed up from ground level, where it broke off, to a height of about six feet; it was apparent that, except for a few inches of sound wood on the south side, all

the base of the trunk was completely rotten, obviously diseased by fungi or bacteria. The rotten wood was soft and crumbly and thoroughly permeated by "black lines".

III. ISOLATION OF THE CAUSAL ORGANISM

By means of a borer, many sterile isolations were taken from the diseased timber in various parts of the stem and roots. The isolations were grown on media known to be favourable to the growth of most fungi and many bacteria, but all isolations produced only remarkably pure cultures of *Ustulina vulgaris*. Eventually the identity of the causal organism with this species of fungus was presumed on the following grounds:

(1) Proximity of undoubted fructifications of Ustulina to the

diseased timber.

(2) Similarity of cultures isolated from the diseased timber with those of standard cultures of *Ustulina*. This diagnosis was confirmed by Mr Cartwright (Head, Mycology Department, Forest Products Research Laboratory, Princes Risborough), both from cultures supplied by the writer as well as from his own independent isolations from the timber.

(3) In spite of repeated trials no other fungus could be isolated.

(4) The details of the disease conditions corresponded very closely with those described, as caused by *Ustulina* on certain tropical trees,

by other investigators.

(5) Artificial infection with the isolated fungus into sound lime wood produced only symptoms of disease identical with those occurring in the naturally infected wood. This was true in inoculation into dead wood as well as in artificial infection into living trees.

(6) The fungus was re-isolated from the artificially infected wood—dead and living—and again pure cultures of *Ustulina* were obtained.

(7) Artificial inoculation into lime with a culture from authentic *Ustulina* spores produced exactly the same symptoms as the above.

The invariable presence of *Ustulina* in the diseased wood does not preclude the possibility of its being a secondary rather than a primary infection, but it is suggested that the evidence produced in the course of this paper tends to disprove this hypothesis.

IV. General observations on the type and extent of the disease

(1) Macroscopic examination

Through the kindness of the Bursar of Magdalen College, I was able to have the whole tree for the purpose of investigation, and both stem and root were cut up into sections.

In the sections of the stem and root the so-called "sound wood"

was the normal wood of the lime, and, structurally, calls for no special comment. It surrounds the discoloured and diseased wood on all sides except that which is completely decayed out to the bark. Cultures of sterile borings from this wood produced no evidence that it contained fungus hyphae.

(a) The stem.

The base of the tree was badly smashed up when it broke off, and the first section which could be cut corresponded to a height of about six feet above ground level; other sections were cut as shown in the following table and will, in future, be referred to by these numbers:

Number of sections ... 1 2 3 4 5 6 Height in feet 6 10 12 14 15 18

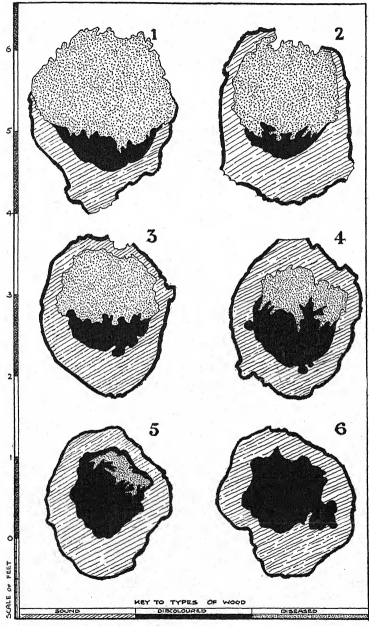
Text-fig. 1, diagrams 1-6, shows diagrammatically the appearance of each of the sections cut at the above levels. The diagrams are from actual tracings of the sections reduced to one-eighth natural size. All the sections are arranged so that the side which originally faced north is uppermost. Plate I, fig. 1, shows a photograph of section 1.

In each of the first five sections, areas representing three distinct types of wood are recognisable; these three types will be referred to as: (i) sound wood; (ii) discoloured wood; (iii) diseased wood.

The sixth section—at eighteen feet—showed only the first two types. All the sections show the relative extent and distribution of the different types of wood in the trunk, and the discoloured and

diseased wood will be discussed separately in greater detail.

(i) The discoloured wood of the stem. This is brownish red and is. therefore, sharply marked off from the diseased wood on the one hand and the sound wood on the other, as both of these are light coloured. The line of demarcation between the discoloured wood and the sound wood is often specially emphasised by a narrow zone of the deeper colour which is found on the outer edge of the former. The successive sectioning of the trunk brought out the fact that the discoloured wood extended to a height of about twenty-four feet, i.e. about eight feet beyond the height actually reached by the diseased wood. In transverse section this discoloured wood shows a tendency to occupy a more or less central position in the stem, simulating heart-wood (lime is a sap-wood tree), and seems to be identical in appearance with the type of discoloration popularly known as "dark-heart", "black heart", "red heart", etc., which may exist as a physiological or pathological condition in all kinds of trees. It is more evident on that side of the diseased wood which is farthest from the (presumed) point of infection; this suggests that (a) it was present in the stem before the disease started, or (b) that it may have been



Text-fig. 1. Transverse sections of the lime stem, cut at different levels (explanation in the text).

produced by the advance mycelium of the fungus and so be a preliminary symptom of decay. In neither longitudinal nor in transverse section does it show any regional relation to the annual rings, but, as its outer edge tends to follow the course of the vessels, in longitudinal section this edge appears as a more or less straight line. In the lowermost sections it has been largely "overrun" by diseased wood and so appears merely as a comparatively narrow zone, and even this may be present at one part of the circumference only (Textfig. 1, diagrams 1-4). In the uppermost sections it forms a relatively wider zone entirely surrounding the diseased wood (diagram 5), until, in the highest section of all, it alone occupies the central region of the stem. It seems, in fact, that the discoloured wood bears a closer relation to the stem than it does to the fungus, for its outline is more or less concentric with the outline of the stem, whereas it shows no definite relation to the outline of the diseased area.

Discoloured wood of the type mentioned above has been noted by Hubert ((11), p. 533) on a species of lime infected by *Pholiota adiposa*, and has also been referred to by Sharples ((19), p. 12) in

connection with *Ustulina* disease of rubber.

Numerous sterile borings from this wood were cultured but gave

no evidence that the wood contained mycelium.

(ii) The diseased wood of the stem. This extends up the trunk from ground level to a height of about sixteen feet. Reference to Text-fig. 1, diagrams 1-5, will show that towards the base of the tree the diseased wood occupies the greater part of the cross-section, but its area gradually decreases from the base upwards till, at a height of eighteen feet, it is no longer present; at the same time it tends to become more centrally placed in the stem, indicating a probable preference for the older tissues or for the discoloured wood. In the transverse direction, the appearance of the diseased area suggests a progressive spread of the mycelium from the wound on the north side, and it is obvious that the disease spreads more rapidly in a longitudinal than in a transverse direction.

The diseased wood is light in weight, crumbly in texture and of a lighter colour than normal lime wood. On the more completely diseased sections—in general on sections nearer the base than the lowest section illustrated in the diagrams—the wood is typically permeated by "black lines". These are irregularly distributed and presumably make their appearance as the wood "dries out" as the result of exposure, produced naturally or artificially. They are found in wood which is on the exposed surface of the trunk but not in the diseased wood which is situated some distance from the outside. On allowing the sections to dry naturally in the laboratory, however, black lines became apparent where they had not been noticed previously. They often extended across the wood more or less parallel

to and a short distance from the cut surface, and a black line was almost invariably produced (after drying out) at the junction of the diseased and the discoloured areas. The lines sometimes enclosed irregularly distributed areas of wood of a dark brown colour.

Though definitely and invariably associated with decay by *Ustulina*, these black lines are not a specifically diagnostic character, as they are present in many kinds of diseased timber, being produced by

some thirty different species of fungi.

The above facts on the appearance of the diseased wood as seen by the naked eye agree in all essential details with those described for *Ustulina zonata* on rubber by Sharples (19) for Malaya, by Steinmann (21) for the Dutch East Indies, and by Weir (25) in the Amazon Valley.

As stated previously, cultures of *Ustulina* were consistently produced from sterile borings of this type of wood, though less readily from the older parts than from the region nearer to the edge of the advancing mycelium.

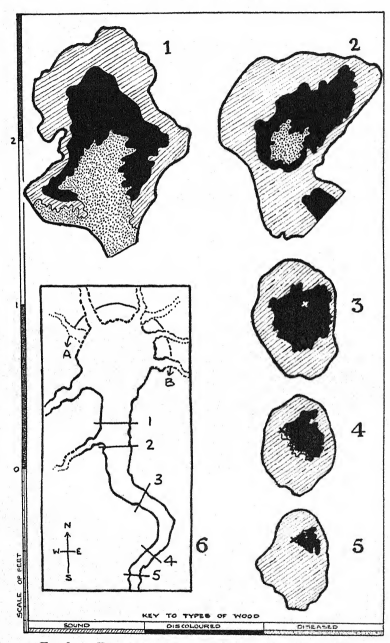
(b) The root.

The root system of the tree was also examined. Text-fig. 2, diagram 6, shows the general lay-out of the roots as they appeared after excavation. There was no tap root, merely a bunch of fibrous roots about three feet deep, the largest being about $2\frac{1}{2}$ inches across. The roots on the north side of the tree—from A to B on the diagram—that is to say the roots arising from below the wound on the stem, were quite rotten and broke when dug up. On the whole the roots were small; there was one large lateral root on the south side and growing towards the south, and a smaller lateral on the south-west. All these roots were infected by *Ustulina*. The largest root (south side) was sectioned as follows:

Number of section ... I 2 3 4 5 Distance from trunk ... 9'' I'3" 2'6'' 4'6'' 5'3''

The root sections showed the same three types of wood as did the stem, and Text-fig. 2, diagrams 1-5, shows the relative extent and distribution of these types as seen in transverse section. As in the stem, the diagrams are from tracings of the timber, but here are reduced to quarter natural size. All the diagrams (except diagram 6) are arranged so that the morphologically upper side of the root is uppermost. Plate I, fig. 2, is a photograph of root section 1. The first two sections show all the types of wood, but on the last three sections, only sound wood and discoloured wood are represented.

(i) The discoloured wood of the root. From the diagrams in Text-fig. 2 it will be seen that this is particularly well represented in the root. It is rather darker in colour than the corresponding area in



Text-fig. 2. Transverse sections of the lime root cut at different levels (explanation in the text).

the stem—this is well shown in Plate I, fig. 2—though here the deeper colour is emphasised rather out of proportion due to the greater moisture content of the root at the time of taking the photograph—the discoloration always being more definite when the wood is damp. This discoloured wood extends down the root for a distance of about five feet six inches, i.e. about three feet beyond the diseased wood. It exhibits the same tendency to be restricted to the central regions that was noticed in the stem and, especially in the upper part of the root (Text-fig. 2, diagrams 1 and 2), its outline follows that of the periphery of the root sufficiently closely to suggest a physiological connection. The fact that there is no relation between the distribution of the discoloured area and the annual rings is well emphasised in the more distal part of the root—as shown in Text-fig. 2, diagrams 3, 4 and 5—where the centre of the root is marked with an X. Though this discoloration sometimes approaches very close to the edge (Text-fig. 2, diagram 3), yet it never extends right out to the bark.

The culturing of sterile isolations from this wood did not produce

any mycelium.

(ii) The diseased wood of the root. There is, of course, some doubt as to the exact position of the original infection, but it would appear that the fungus travels less quickly in the root than in the stem in both the longitudinal and the transverse direction. There was no evidence of any diseased wood at two feet from the junction with the trunk, and the area of this type of wood in the cross-section of the most diseased part is comparatively small. The diseased area is found on the lowermost side of the root and, as in the stem, it spreads towards the centre. In all structural details the diseased wood of the root is similar to that of the stem. Black lines, which are not evident in the freshly cut timber, make their appearance as the wood dries out.

Sterile borings from this wood consistently produced cultures of *Ustulina*.

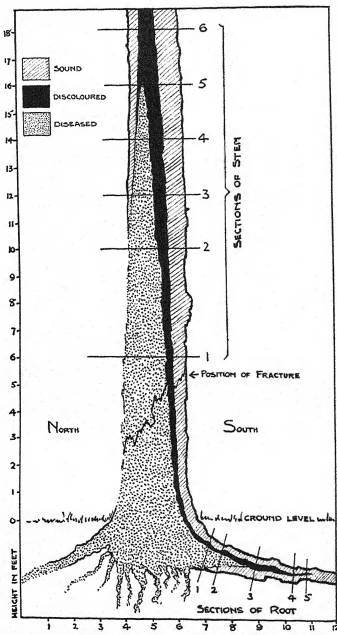
(c) Distribution of disease in the tree.

From the successive transverse sections of the stem and root and from the records obtained when these are cut longitudinally, it is possible to reconstruct the general distribution of the diseased tissues in the tree as a whole. Such a reconstruction is illustrated diagrammatically in Text-fig. 3, which is self-explanatory. This is a typical example of the effect of *Ustulina* as a wood-destroying fungus.

(d) Superficial mycelium.

After the sections had been cut for about a week, a dense mycelial growth appeared on the surface of certain sections of both stem and root. It grew out of the diseased wood and was never found on any

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Text-fig. 3. Reconstruction of a longitudinal section through the tree (explanation in the text).

other type of wood; the outline of this mycelium followed the outline of the diseased wood exactly. This superficial growth showed first and grew most luxuriantly on the most distal sections of stem and root, i.e. the mycelium seems to be most active on the growing margin. This was confirmed by observation of the transverse sections, for the fungus always appeared first on the extreme edge of the diseased wood where it bordered on the discoloured wood, the part farther from the discoloured wood showing a much more scanty growth. At a later date the mycelium was to be found on the lower sections but—in the stem—it became progressively less vigorous towards the base of the trunk, that is to say on the more completely diseased wood. Plate I, fig. 3, shows a "close up" of this superficial mycelium as it appeared on the uppermost root section. This is a typical example of the appearance of Ustulina mycelium as seen in nature, and this is, moreover, almost identical with its cultural aspect.

This type of superficial mycelium is figured by Schrenk (18), Plate V) in connection with a disease of ash caused by *Polyporus fraxinophilus*, and also by Heald (19), Fig. 246, p. 784), who shows a surface growth of the mycelium of *Stereum purpureum* on cross-section

of apple timber.

(2) Microscopic examination

Before going on to the more detailed examination it may be well to state certain assumptions which will tend to explain the course of the investigation. In a section of the diseased trunk, e.g. section 2 in Text-fig. 1, it is assumed that:

(a) Infection took place at a point in the region of the wound on

the north (uppermost) side.

(b) The spread of the mycelium from this point was fan-wise in the general direction of the opposite side of the stem, that is through the discoloured wood if that was already present, or converting the sound wood into discoloured wood as it penetrated it. In the latter, one is assuming discoloration to be the initial stage of decay.

(c) The actual decay, which proceeded more slowly, in the same sense, then gradually converted the discoloured wood into diseased

wood, i.e. disintegration is the ultimate stage of decay.

With the idea of examining the timber from this point of view a strip about four inches wide was cut down the centre of stem-section 2. Plate I, fig. 4, shows a photograph of this strip in transverse and in radial section. The strip is arranged so that the sound wood is uppermost (the examination being carried out from the periphery inwards), and all the subsequent drawings have the same orientation.

From this strip, pieces of wood were sectioned, either by hand or by means of a wood-cutting microtome, and both before and after

staining were examined and drawn.

Many stains were tried in the course of the microscopic work, but eventually it was decided that the most useful were iodine, chlorzinc-iodine, Mäule's (14) and Cartwright's (5). The most generally satisfactory was the last named because, besides being permanent and demonstrating the fungal hyphae very distinctly, it had the additional advantage of tending to differentiate—using Mäule's stain as a criterion of comparison—between cell walls which gave a lignin reaction and those which gave a cellulose reaction, though it must be admitted that considerable experience in the use of the stain was necessary before one could be satisfied on this point. The unreliability of lignin determination by staining methods is well known, hence, in all cases of doubt Mäule's stain was used, because in the opinion of investigators such as Crocker(6), Harlow(7) and Phillips(17), etc., it is the most satisfactory. The microscope drawings were done with the aid of a Leitz projection apparatus.

In writing up this part of the work, the matter was considered under the respective headings "The effect of the fungus on the timber" and "The effect of the timber on the fungus", though con-

siderable overlapping was unavoidable.

(a) Effect of the fungus on the timber.

(i) The discoloured wood. The sound wood hardly comes into this at all, so it will be well to start with the discoloured wood (see Plate I, fig. 4, B, C). This is about 1 cm. wide and is seen to consist of two regions:

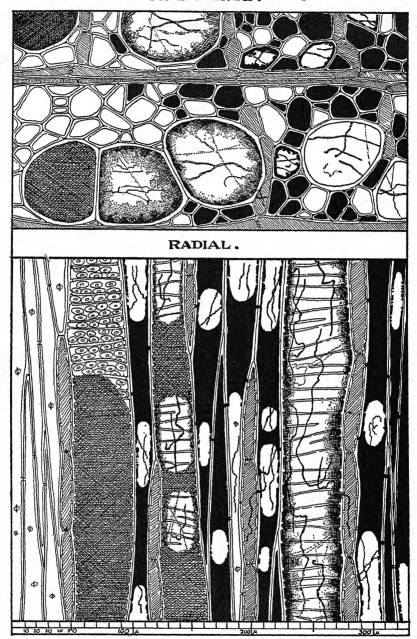
(1) A peripheral zone about 5 mm. wide, dark red-brown and bordering on the sound wood. This type of wood will be referred

to subsequently as "D2".

(2) The discoloured wood itself which, being very variable in width, can hardly be described as a "zone". This is of the same general colour as the D2 but less deep in tone; it will be referred to as "D1".

In both, the discoloration is largely consequent on the deep colour of the cell walls, those of the rays and wood parenchyma being yellow while those of the vessels and tracheids are red-brown. In addition to the deep colour of the walls, however, the D2 wood shows the phenomenon of occlusion of both vessels and tracheids by dark brown infiltrations. These have the superficial appearance and give the same staining reactions as the substances described by certain previous workers under the somewhat indeterminate heading of "wound gum". An interesting feature of these products is that when stained with Cartwright's stain, the products in the vessels stain blue (? cellulose complex) while those in the tracheids stain red (? lignin complex).

In the outermost half of the D2 region only the vessels are filled with these products, but in the innermost half the tracheids also are



Text-fig. 4. Transverse and radial sections through the timber in the wound-gum region."The red-stained infiltration products in the tracheids are in full black, while the blue-stained products in vessels are cross-hatched.

filled; in fact, the massing of the products in this region is so abundant that practically every tracheid is filled with them. The products, though commonly found in vessels and tracheids, are comparatively infrequent in rays and parenchyma. Text-fig. 4 shows the transverse and radial appearance of this region of the wood, the blue-staining products in the vessels are cross-hatched, while the red-staining products in the tracheids are in full black. From the radial section it will be seen that the products are not continuous throughout the whole length of the tracheid but are interrupted at irregular intervals by spaces. This produces in transverse section the erroneous effect that a considerable number of the tracheids is empty. In certain vessels products are found to be adhering round the walls, while the centre is devoid of such products; these vessels invariably contain hyphae, and it is suggested that the products are probably being digested by the fungus.

The Dr wood, though having the coloured walls, does not show the filling up of the cells to any extent; occasionally cells are so filled

but it is comparatively rare.

The discoloured wood of the root is essentially similar to that of the stem, except that there is less evident distinction between the D_I and D₂ types of wood owing to the fact that all the cells of the D_I region of the root show a higher proportion of infiltration.

Though this discoloration may be taken to represent "incipient decay" it must, from the decay point of view, be taken as an indication rather than a fact, as the most careful examination failed to reveal any structural disintegration of the timber in the discoloured

region.

(ii) The diseased wood. The diseased wood is delimited from the discoloured wood by the black line, as shown in Plate I, fig. 4, at C. Behind this line the wood is light in colour, crumbly in texture, and has, in fact, all the superficial characters of diseased wood. Microscopic examination, however, shows that disintegration is not a direct function of the black line itself, for actual disintegration commences about 5 mm. behind the black line. This is in agreement with the observations of Hiley ((10), p. 156) on the black line of Armillaria mellea, where he says "when looked at with the microscope it is remarkable how little difference can be seen in the wood on the two sides of the black line...nevertheless at some distance behind the black line marked delignification does take place". Besides the black line which is found on the edge of the diseased wood, other apparently similar lines occur scattered indiscriminately throughout the older parts of the diseased wood; as these black lines consist entirely of hyphae they will be discussed later.

The diseased wood some few millimetres behind the black line shows the first stage of decay. This decay is always more marked on the autumn than on the spring-wood side of the annual rings. The large vessels of the spring wood appear to be unaffected; they remain intact, show no change of structure and no difference in their reaction to stains. The tracheids of the spring wood, however, are beginning to show some sign of change, their walls are thinner and less rigid than those of the sound wood, though they continue to give a lignin reaction. Below the ring, decay is more advanced; the walls of the vessels are still unaffected, but the tracheid walls have become much thinner, have lost their rigid structure and appear wavy and fragile; often there seems to be little but middle lamella left. They show evidence of considerable delignification.

In both spring and autumn wood, the rays and wood parenchyma

remain apparently quite unaffected.

In very badly diseased timber, farther back from the line, the above state of affairs is emphasised; vessels, rays and wood parenchyma still show no sign of disintegration, but the tracheids of the spring wood have now reached the stage described above for the autumn tracheids, while the tracheids of the autumn wood here are completely disorganised, the walls have broken down completely and only a few scattered fragments of the middle lamella are left.

By the use of suitable stains, the course of the disintegration can be suggested. Both Mäule's and Cartwright's show when the wood fails to give a lignin reaction—the former by the absence of colour and the latter by the presence of a blue colour. It is difficult to state that the absence of the lignin reaction indicates that the delignified wall was reduced to cellulose as no cellulose indicators (such as chlor-zinc-iodine) gave a consistently positive result. The experimental use of the stains on other timbers, however, tended to indicate the probability that it was so, and this has been tentatively assumed.

Stages in the disintegration of the tracheids can be said to occur

as follows:

The first stage is the appearance of delignified spots in the wall. With Mäule's stain the effect is as if a piece of the wall has been "bitten out", but Cartwright's stain shows that there is still wall substance in these spots, as, with this, they stain blue. This sort of thing develops until large parts of the wall become delignified, and often it is only at the corners that any sign of lignification remains. Eventually the whole of the wall becomes delignified and at the same time seems to be much thinner, *i.e.* it appears to collapse simultaneously with the disappearance of the lignin. At this stage the wall may lose its rigidity and become wavy or even broken. The last stage of wall disintegration is that only the middle lamella is left and finally the tracheids disappear entirely leaving an empty space surrounded by the non-disintegrated elements.

The skeleton formed by the vessels, rays and wood parenchyma

when the tracheids have disappeared, ensures that lime wood decayed by *Ustulina* always retains a certain stability even when the specific gravity is reduced to about 0.2 and the wood crumbles readily in the fingers. An example of the skeleton effect produced in a very badly decayed piece of timber is illustrated in Text-fig. 5. This also illustrates the relative frequency of fungal hyphae at this stage of decay.

(b) Distribution of the fungus mycelium in the timber.

The mycelium penetrates the whole of the diseased and the discoloured areas. In spite of the fact that no cultures were obtained from borings taken from the discoloured wood, microscopic examination reveals the presence of hyphae in that wood. Taking the strip of wood (Plate I) in the same order as before, it is found that hyphae are first seen in the sound wood outside the discoloured area. For about 5 mm. beyond the outer margin of the discoloured wood, the hyphae are fairly abundant—though not as numerous as in the discoloured wood itself; and some—relatively few—hyphae are found to extend as far as 10 mm. beyond the margin. Unless otherwise stated, all the hyphae are narrow—about 1 μ in diameter.

In the D2 zone the hyphae are fairly abundant; there are approximately three hyphae in each vessel and they are rather less frequent in the tracheids. It has been stated before that in the outer part of D2 it was only the vessels that showed decomposition products, and this would appear to be correlated with the distribution of the hyphae. In the innermost part of this region, where the tracheids also showed infiltration, hyphae are more numerous in the tracheids, but where the infiltration is very dense they are either absent or indistinguishable. It was noticed that all vessels which had the products only

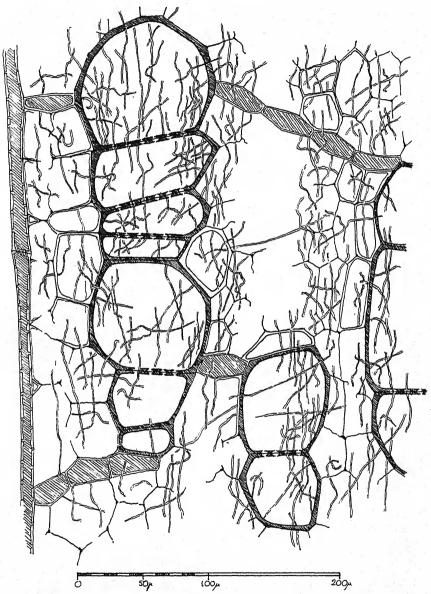
round the margin of the cell invariably contained hyphae.

The D_I region contained relatively fewer hyphae; approximately half the vessels and rather fewer of the tracheids contain about one or two hyphae each. It would appear, therefore, that there is a slightly increased development of hyphae in the outer region of the D₂ margin of the discoloured wood. In both regions hyphae were

occasionally found in the rays and parenchyma.

The next region is the black line. It is usually a few cells wide, and the cells comprising it are densely filled with black contents so that the whole line appears as an amorphous mass (Brooks(2), p. 160). In thin sections, however, and particularly on the edge of the line, the "tylose" origin of these lines, as commented on by several investigators, is very obvious. This appearance is illustrated by Small ((20), Plate II, fig. 9), by Hiley ((10), pp. 155 and 157) in connection with Armillaria on larch, and by Campbell ((3), Plate III, figs. 1, 2 and 3).

In the black line no hyphae of the ordinary type could be dis-



Text-fig. 5. A transverse section of a very diseased part of the timber (explanation in the text).

tinguished owing to the dense massing of the black substance. On the side of the line which is towards the discoloured wood, however, large septate hyphac—about 4μ in diameter—appear to grow out from the black line substance, and they extend vertically up the vessels, tracheids and parenchyma for a distance of about 0.5 mm. and for approximately the same distance along the rays. These hyphae are all filled with a dark-coloured substance in the part which is nearest the line, but this discoloration gradually decreases towards the more distal parts, and the ends of the hyphae are hyaline. This state of affairs—which does not appear to have been mentioned by previous investigators—is illustrated in Text-fig. 6 which shows the transverse appearance, and particularly well in Text-fig. 7 which shows the radial appearance of the timber.

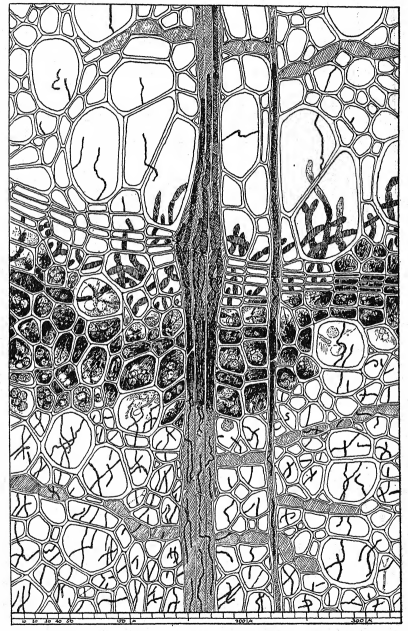
In the diseased wood behind the line, hyphae are present in large numbers; practically all the cells contain them, and sometimes the cells are literally packed with hyphae. These are all of the narrow type. Even here decay is, to some extent, localised; there may be regions where decay is slight with comparatively few hyphae, and other regions not necessarily nearer the original infection—where decay is very advanced and the hyphae extremely numerous. This type of vigorous decay may occur over a region several inches deep, but in the much older parts, where the wood is completely rotten, and has been so for some considerable time, hyphae are relatively infrequent.

Black lines also occur indiscriminately throughout the diseased wood, but, in lines which are not situated at the growing margin of the mycelium, *i.e.* at the edge where the diseased wood borders on the discoloured wood—the line merely shows the usual characteristic appearance, and there is no evidence of the above-mentioned large hyphae. Small hyphae, also, are often much less numerous in the cells associated with the black lines which are situated in the "older" parts of the decayed timber.

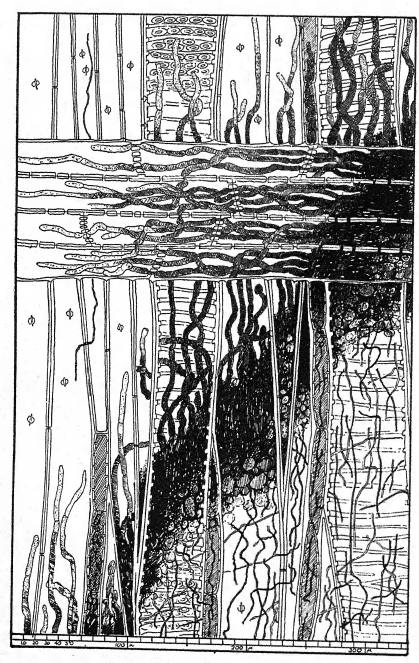
Previous workers have suggested that the black line is intimately connected with the process of timber decay, but, up to the present, no adequate account of the significance of the line in this connection has been produced, and I am carrying out investigations on this subject.

(c) Hyphal penetration.

In general, penetration always seems to be by means of the pits; the presence of bore holes was never detected. Nutman(15) found that, in wood subjected to the action of *Polyporus hispidus* for four months, penetration was always by means of pits; and that later penetration of vessels, rays and wood parenchyma (except in the wall bordering on the fibres) was also by means of the pits, but the fibres showed extensive penetration by bore holes.

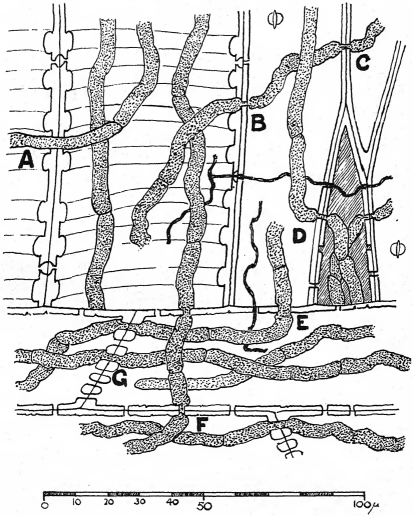


Text-fig. 6. A transverse section of the junction between the diseased and the discoloured wood, showing the black line and the relative distribution of the hyphae.



Text-fig. 7. A longitudinal section through the same region as shown in Fig. 6.

The $I\mu$ hyphae were about the same size as the pits and went straight through without apparent alteration. The larger hyphae, on the other hand, were sometimes constricted when they passed



Text-fig. 8. Examples of hyphal penetration (explanation in the text).

through a pit and sometimes not, according to circumstances. Text-fig. 8 shows examples of the various types of penetration.

It will be seen that when the larger hyphae pass from vessel to vessel by means of the bordered pits, the border seems to be dissolved

away and, the pit then being about the same size as the hypha, the latter passes through without constriction (A). When passing from vessel to tracheid or from vessel to parenchyma the hypha is not constricted when it passes through the vessel wall, but is constricted when passing through the wall of the other element (B). In passing from tracheid to tracheid or from tracheid to parenchyma, definite constriction takes place (G and D). When hyphae are passing through the side walls of the rays the degree of constriction varies with the size of the pits, which themselves vary in diameter (E and F), but when passing through the end walls of the ray cells where the pits are rather large, the constriction is slight (G).

Usually the hyphae, having been constricted when passing through a pit, seem to swell up to their original size on the other side, but in certain comparatively rare instances this is not so, and the hyphae continue on the other side for some considerable distance as "narrow" hyphae. Rarely the hyphae flatten out into a disc before passing through a pit; here the actual penetration is probably of the "pegoutgrowth" type, though this has not been seen. In transverse section, the characteristic features of penetration appear essentially the same

but are less easily discernible.

V. Conclusions

From the foregoing it seems reasonable to conclude that *Ustulina vulgaris* is capable of forming a very definite disease of standing lime, and from the evidence it is suggested that this fungus is to be regarded as a wood destroyer producing a white rot of the timber to such an extent that the tree may be completely destroyed and the timber reduced to a state where it has little, if any, commercial value. Speaking generally, the disease can be classified as a "white rot" and appears to belong to that type of white rot which falls into the Group II of Campbell(4), *i.e.* a white rot "in which the cellulose with its associated pentosans is attacked in the early stages and in which the incidence of the attack on lignin and the pentosans not in cellulose is delayed".

This paper deals only with the observational facts of *Ustulina* disease of a certain lime, the results of inoculation experiments and experimental work generally not yet being complete are postponed to a subsequent paper. At the moment, however, it is possible to state that all the evidence up to now tends to confirm and justify the opinion that *Ustulina* must be considered as possessing the poten-

tialities of a pathogen of economic importance.

VI. SUMMARY

1. A diseased lime was investigated with the object of determining the cause of disease. The only fructifications present were those of Ustulina vulgaris Tul.

2. The causal organism was isolated and proved to be Ustulina;

this was confirmed by reinfection and subsequent isolation.

3. The tree was cut into sections and the extent of the disease in the stem and root was established.

4. Microscopic examination showed that the fungus attacked the cell walls of the tracheids, leaving the vessels, rays and parenchyma

practically unaffected.

5. The distribution of the fungus mycelium in the timber was determined by microscopic examination and the types of fungal penetration described.

6. It was concluded that Ustulina causes a white rot of lime.

I am indebted to my Research Assistant, Miss E. M. Ellis, B.A., B.Sc., for valuable help in connection with this work.

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EXPLANATION OF PLATE I

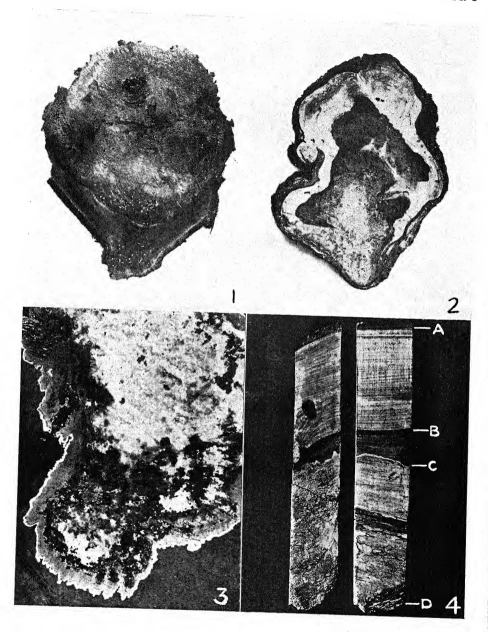
Fig. 1. Transverse appearance of the lowest section of the stem, cut at a height of six feet from ground level.

Fig. 2. Transverse appearance of the first root section, cut at a distance of nine inches

from the junction with the trunk.

Fig. 3. A "close-up" view of the superficial mycelium which appeared on the above root section.

Fig. 4. The transverse (left) and radial (right) appearance of a strip of timber cut down the centre of stem section 2 (explanation in the text).





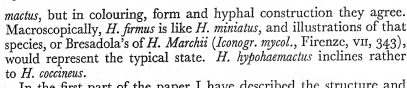
HYGROPHORUS WITH DIMORPHOUS BASIDIOSPORES

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(With 8 Text-figures)

I have found in Malaya two species of Hygrophorus with dimorphous basidiospores. Side by side, on the same fruit body, occur large spores on large basidia and small spores on small basidia. The large spores have dense contents and roughly twice the linear measure of the small spores which are hyaline and vacuolate and often with a slightly different shape. The meaning of this dimorphism, which is unknown in any other Basidiomycete, cannot be as great as one would at first suspect and is by no means equivalent to true cryptogamic heterospory. For, in the first place, both species are very similar in all other respects to certain common and well-known European members of this large, unspecialised genus, and unlikely, therefore, to have evolved a new physiological condition. And, in the second place, both kinds of spore invariably occur on the same fruit body, on adjacent hyphae, and all the basidia are tetrasporous and clamped (the small basidia rarely being disporous), so that a sexual or karyokinetic difference can scarcely exist. It is perhaps just the full consequence of a vegetative dimorphism which occurs among the hyphae of the fruit body of these, as in many other agarics: in the primordium, all the hyphae are similar, but some inflate strongly and some hardly at all, and so of the basidial branches, which are initially similar, some inflate strongly to produce the large spores and others but little to produce the small spores. But I have examined only the morphological aspect.

One species, Hygrophorus firmus, is known already from Ceylon, where Petch remarked on the dimorphism(4). In Malaya, it varies exceedingly in shape, colour, size and spore characters, surpassing even Laccaria laccata or Omphalia umbellifera, and, reckoned with its varieties, it is one of our commonest toadstools. The other, which I propose to call Hygrophorus hypohaemactus from the rich colour of the hypodermal tissue, is new and probably rare. They differ chiefly in a way which is not infrequent in the genus. In H. hypohaemactus the hyphal ends on the surface of the fruit body have mucilaginous walls and their cells are not inflated, whereas in H. firmus they have firm, not diffluent, walls and at least over the pileus they are inflated. There is also less dimorphism in the spores and basidia of H. hypohae-



In the first part of the paper I have described the structure and development of the fruit body, which offers certain points of interest in the origin of the pileus, the marginal growth of the limb and gills, and the expansion mechanism of the fruit body. I have also included a summary of the bionomical observations which I have made in Singapore on the fruit body of *H. firmus* var. *stratiotes*: they will be published in detail at a later date together with illustrations of the development and shape of the fruit body in both species and varieties.

The structure of the mature fruit body of Hygrophorus firmus var. stratiotes

The stem. The hyphae are very compact and chiefly of two kinds: either they are strictly longitudinal with inflated cells, 120–1800 \times 10–35 μ , or narrow and more or less interwoven with the cells passively extended, 20–200 \times 3–5 μ . Both kinds are clamped and the narrow ones are more frequently branched, the branches arising from any part of the cell, generally from the distal half. The contents of the wide hyphae are clear and vacuolate, though sometimes vitreous at the tapered ends of the cells; those of the narrow hyphae are vacuolate and rather smeary-looking, or they are frequently oily throughout, but initially, in the primordium, all the hyphae are similar and the narrow ones begin to acquire their characteristic appearance when the fruit body is about half-developed.

Many intermediates occur between the two kinds, and the one often gives rise to the other. Generally the cells of a hypha are inflated throughout its length or not inflated, but often a few inflated cells may lie between rows of narrow cells, and even the same cell may be inflated at one end only or in the middle, or occasionally

in several places, so appearing nodose.

At the surface of the stem the hyphae are all narrow, $3-8\,\mu$, and scarcely inflated. Some of the hyphal ends project obliquely up to 200 μ as narrow filaments with one or two septa and occasionally branched, but they are scattered and there is no palisade of caulocystidia: the terminal and subterminal cells of the hyphae may be slightly inflated, $25-45\times5-14\,\mu$, especially near the apex of the stem. The hymenium begins abruptly with only a few small tufts of cystidia and sterile basidia preceding it.

The hyphae bordering the hollow of the stem are mostly narrow

and not inflated.

The stem is waxy and brittle, owing to this arrangement of the

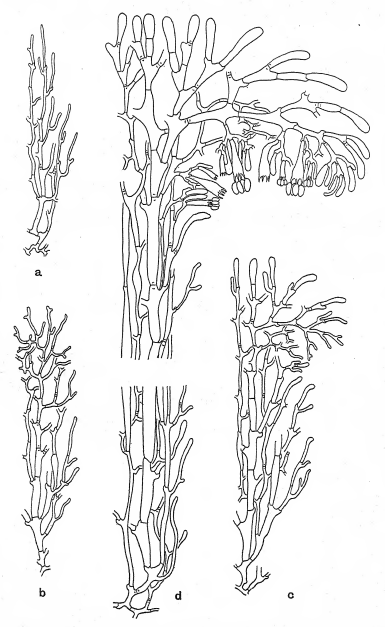


Fig. 1. Diagrams of characteristic hyphae from successive stages in the development of the fruit body of *H. firmus*: a, the primordial shaft; b, the primordial shaft on development of the pileus; c, with primordial limb and incipient hymenium; d, the mature fruit body.

hyphae, and snaps easily under the weight of the pileus when dis-

placed from the vertical.

THE PILEUS. The structure is like that of the stem, but the inflated hyphae are frequently 40 μ wide and rarely so long (up to 600 μ). At the surface the hyphae are all inflated. Over the centre their ends form an irregular pile of simple obtuse filaments, two to four cells long, and projecting freely or in groups up to 300 μ high, which gives the scurfy appearance to the disc: their cells are $60-200 \times 9-25 \mu$, and rarely branched. Over the limb the hyphal ends are radiating, adpressed and rather narrower so that the pile is scarcely recognisable and the surface is inoderm, while near the margin their cells are only 20-60 \times 7-15 μ and pass conformably into the hymenium.

THE TRAMA. The structure is like that of the pileus, but most of the hyphae are inflated with shorter cells, up to 200 μ long, but as wide as in the pileus. The trama and the tissue of the pileus immediately above the gills are hygrophanous with water, or very

dilute mucilage, between the hyphae.

THE SUBHYMENIUM. This tissue is also hygrophanous. It is loosely plectenchymatous, being composed of narrow hyphae with cells $8-35 \times 2-4 \mu$, not inflated, but pulled apart through the intercalary

growth of the hymenium.

THE HYMENIUM. The basidia are dimorphic. The large basidia arise deeply in the subhymenium and terminate some distance beyond the general level of the small basidia; the large spores project well beyond the small spores. Both kinds of basidium are borne on the same hyphae, the large basidia being derived from deep-seated laterals and the small basidia from the superficial laterals; intermediates are rare. The small basidia far outnumber the large basidia which are dispersed evenly without bunching, though they are generally absent from the first-formed parts of the hymenium at the junction of the gills with the stem. As the young pileus is expanding, the basidia mature acropetally from the stem apex to the margin of the limb and obliquely outward down the gill to its edge, but very soon the gills become aequihymeniiferous, and large and small spores are shed together. There are no pleurocystidia.

THE GILL EDGE. The construction varies but the edge is always sterile. The simplest condition shows a narrow strip of cheilocystidia, formed from the modified ends of the down-growing tramal hyphae. Transitions between the cheilocystidia and small basidia are frequent, as clavate cells with abortive sterigmata and even rudimentary spores. More complicated conditions show the ends of the tramal hyphae, $3-5 \mu$ wide, far exceeding the hymenium and matted in a loose plectenchyma, while their ends may inflate, 6-10 µ, and appear in

surface view as decumbent or immersed cystidia.

THE COLOUR. As is general in Hygrophorus, the brilliant red or yellow

(and its modifications in the other varieties of H. firmus) is due to a pigment dissolved in the cell sap of the superficial hyphae. The internal hyphae and those of the hymenium are colourless.

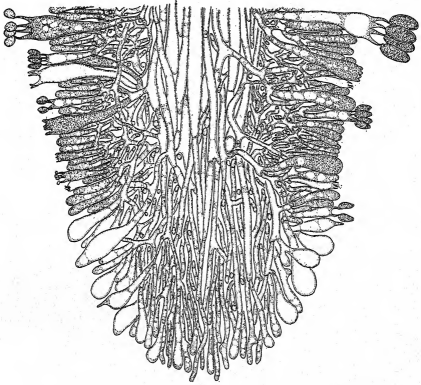


Fig. 2. The edge of a gill of *H. firmus* on cessation of downgrowth; $\times 400$.

THE DEVELOPMENT OF THE FRUIT BODY OF HYGROPHORUS FIRMUS VAR. STRATIOTES

Macroscopic features. Development is gymnocarpic and direct, with exogenous pileus. A conical primordial shaft, $0.5-3.5 \times 1-2$ mm., is first developed; it is initially colourless but soon becomes pale orange save at the tip. A small swelling, 1-2.5 mm. wide, then forms at the tip through outgrowth on all sides. Presently the outgrowth increases in an equatorial belt and the limb develops. Through epinasty the limb becomes convex with the margin slightly incurved. All the while, the part below the swelling is enlarging, elongating and raising the pileus, to become the stem; it soon colours scarlet, though not as deeply as the pileus. The primary gills develop almost as soon as

the pileus, and the secondaries and higher ranks are intercalated regularly at the margin. The pileus begins to expand before the stem is fully elongated. The limb straightens, becomes horizontal and finally is thrust upward by the expansion of the gills which renders the pileus concave or infundibuliform. The stem soon becomes hollow because the internal tissue is disrupted on expansion of the peripheral, and the excessive growth of the gills may disrupt the central tissue of the pileus and cause the fruit body to be pervious to the base.

Microscopic development

The primordial shaft. Composed of longitudinal interwoven hyphae, the primordial shaft enlarges by apical growth and acropetal inflation of the cells. The hyphal tips, $1.5-3 \mu$ wide (mostly $2-2.5 \mu$), are rounded and obtuse with the apical cells 12-55 µ long. They grow monopodially and often branch by lobing from the apical cell without concomitant septation, or from the subterminal cells, which are mostly 12-30 μ long on delimitation. The direction of growth varies so that the hyphae are interwoven, but as a whole the apex of the primordium grows away from the substratum. The enlargement of the cells begins at the base of the primordium, which is therefore conical; when the primordium is barely 0.5 mm. high, the hyphae at the base are beginning to inflate, and when o 5-1 mm. high, the cells in the basal third measure 20-60 \times 4-10 μ . The hyphae on the outside of the growing point of the primordium gradually cease growing: they are left behind on the surface and their ends either remain unmodified, adpressed or slightly projecting, or they may inflate slightly, 5-14 μ wide, as already described: but they are never abundant and do not branch regularly or profusely so that neither a pile nor a palisade is constructed. The tissue of the primordium is hygrophanous throughout, water filling the interstices between the hyphae and rendering the whole translucent.

This is the general organisation of the primordial shaft in the higher fungi, but it has not previously been described in detail for

a gymnocarpic agaric.

The pileus initial. After a short interval, 36-60 hours, from the origin of the primordium, when it is 1-3 mm. long, apical growth is checked. The terminal cells of the longitudinal hyphae at the tip of the primordium begin to inflate, becoming narrowly clavate, 3-5 μ wide. From their subterminal cells numerous laterals arise and, growing to the surface by devious routes from various depths, are checked and branch again in their turn. The apex of the primordial shaft thus swells into a small head composed of narrow, 1.5-3 μ , densely interwoven hyphae, the ends of which are excrescent from the whole surface except over the original apex of the shaft where a dead space has formed. In this dead space the hyphal ends arrange

themselves in a pile which, together with the underlying interwoven tissue, will become the disc, or centre, of the pileus. The dead space extends centrifugally, and simultaneously, the outgrowth from the lower side of the "head" is checked in a zone round the stem apex, and this zone spreads upward over the side of the "head", except along certain radial paths which become the paths of outgrowth of the primary gills. Outgrowth from the head of the primordium is thus checked from above downward and below upward but continues in an equatorial zone as the marginal growing region of the pileus. The form factors begin to play, as usual, in a region where

an active and regular outgrowth has been established.

The marginal growth of the pileus. As the hyphal ends grow out in an equatorial zone round the head of the primordial shaft, they build up the limb. This zone becomes the margin of the limb, the growth of which is monopodial like that of the primordial shaft, but the tissue of the limb, which it constructs, is dorsiventral and the margin becomes incurved. This epinasty is due partly to the dorsiventral structure and partly to a directive growth of the hyphal tips. One can consider the limb as a corticated, multifilamentous, centrifugal and fan-shaped soma, the cortex on the upper side being the sterile pile, that on the lower side being the hymenium, and the wide intervening medulla forming the flesh. The hyphal ends in the marginal growing region behave like those at the apex of the primordial shaft, but those which drop behind are modified according as they lie on the upper, outer or lower, inner side (Fig. 3). If they lie to the outside, their terminal cells enlarge, becoming clavate, and eventually inflated up to 25 μ wide: they become coloured, and branch sparingly from the coloured subterminal cells, and form the pile on the pileus: their proximal parts contribute to the medulla of the limb along with the subterminal cells of the main longitudinal hyphae from the margin. If they lie to the inside, their terminal cells enlarge into basidia and are unpigmented. The terminal cells of the upper sterile cortex enlarge sooner and to a greater extent than the first formed basidia in the hymenium (which are undergoing meiosis), and thus the margin of the limb is pressed down and in, toward the stem, as a mechanical effect of the growth of the pile. But the hyphal tips at the margin tend to grow toward the stem by a curvature independent of tissue pressures and determined by some intrinsic property of the apical cells (Fig. 3). There is no evidence that the hyphal tips are geotropic, although they may be negatively phototropic.

The pileus. As already explained, a close pile is built up over the centre of the pileus from the ends of the longitudinal hyphae of the primordial shaft and their laterals. A similar pile is built over the limb, but as the limb extends, the hyphae of the pile do not inflate or branch as much, but are more or less decumbent. The hyphae



of the pile are deeply pigmented and the unexpanded or partly expanded pileus appears like scarlet velvet. As the medulla extends the pile is disrupted into scurfy particles, becoming almost unrecognisable in mature pilei, and the colour is diluted and gradually fades. In some specimens the decumbent hyphae on the upper side of the limb scarcely branch at all and the limb appears inoderm without

a pile.

The gills. As in Collybia apalosarca (2), the gills develop as ridges along radial paths on the under side of the limb where the outgrowth from the primordial head is not checked. Along these paths, which extend radially with the growth of the limb, the hyphae grow monopodially, as they do at the margin of the limb. The gills are thus formed by the outgrowth of hyphae. This is clear when the tramal hyphae are excrescent to form a loose plectenchyma along the gill edge (Fig. 2). And as the primary gills diverge centrifugally, keeping a constant width, so the secondaries and, in their turn, the higher ranks are intercalated. At the sides of the gill edges the apical growth of the hyphae slows down; their apices bulge out unilaterally at right angles to the tramal hyphae and develop into basidia. The first basidium is generally large. Laterals arise from the subterminal cells and, growing up in the same direction, generally terminate in large basidia: laterals arise from their subterminal cells, and growing up to the level of the large basidia cut off one or two short, subhymenial cells and terminate in small basidia. Sympodial branching then continues profusely, the deep-seated laterals generally becoming large basidia, the more superficial ones small basidia. Thus the level of the hymenium is built out into the gill spaces on a relatively thick subhymenium, and one finds the large basidia scattered among the crowds of small basidia. The large basidia originate in the same way as the pleurocystidia of C. apalosarca, and their stalks traverse the subhymenium. On cessation of growth the hyphal ends along the gill edge enlarge into the cheilocystidia.

The basidia and spores. On account of their size the large basidia are convenient objects for study (Fig. 4). One can distinguish three main processes: the accumulation of cytoplasm in the basidium, the vacuolation of the basidium driving the cytoplasm into the sterigmata and spores, and, concurrently, the stretching and stiffening of the cell wall fixes the shape. The basidium is full sized before the sterigmata develop, and the sterigmata before the spores appear; the maximum of the grand period comes toward the end of the first stage at the apex of the basidium with a secondary maximum in the spores.

The basidia enlarge through the accumulation of cytoplasm. While they are slender hyphal ends, they are finely vacuolate and transparent but, as they grow, numerous anabolites are precipitated as reserve substances to be lodged in the spores, and when they are full-sized

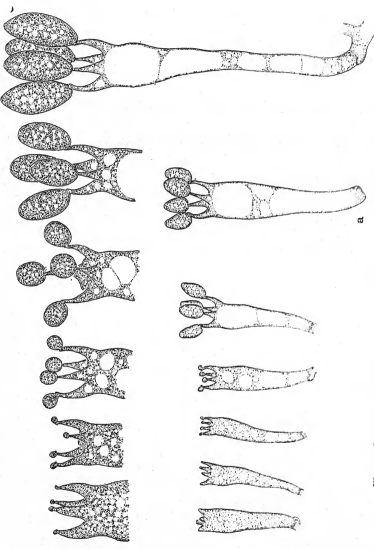


Fig. 4. Stages in the development of large and small basidia of H, firmus var. stratiotes; a, an intermediate basidium; \times 1000.

and cut off by a septum their contents throughout are densely granular and guttate. The wall of the basidium stiffens acropetally leaving four thin spots at the apex, and this leads to the second stage. Vacuoles appear at the base of the basidium as a few small spaces in the cytoplasm, then in the middle of the basidium, and as they enlarge they press upon the denser cytoplasm, the pressure being transmitted to the four weak spots at the apex, and the sterigmata are pushed out as blunt processes: the internal pressure of the basidium now becomes a simple turgor pressure. The walls of the sterigmata are at first elastic throughout but soon begin to stiffen acropetally and the sterigmata taper to an acute apex: this begins the third stage. The spores arise as minute lateral swellings on the abaxial side of the sterigmata just below their tips, at a point where the wall has not yet stiffened. The vacuoles in the basidium continue to enlarge, forcing the granular cytoplasm into the spore rudiments which swell into spherical bodies, indicating that the spore wall is growing uniformly. The spore wall then begins to stiffen acropetally and the distal end of the spore is forced into a blunt cone, like the sterigma. The basidium and its appendages can then enlarge no further although the turgor pressure continues to rise. This instability leads to the discharge of the spores, which has been fully investigated by Buller(1). The cell wall, in a state of increasing tension, finally gives way at the weakest point, at the necks of the sterigmata, and the basidiospores are violently shot off with a drop of escaped fluid.

How the cytoplasm is forced into the spores can readily be gathered from Fig. 4. The vacuoles must be special structures, for there is no general hydration of the cell contents and their walls must be strong enough to press upon the non-vacuolated cytoplasm. The shapes of the basidium, sterigmata and spores are clearly determined by the properties of the cell wall, especially the extent to which it can be stretched locally. Those parts of the wall where the sterigmata and spores emerge, as well as that which finally gives way, may of course be weakened after stiffening, but it is simpler to construe them as direct steps in the continuous development of the organ: the problem is why the sterigmata and spores are blown out only in certain

places.

It is not obvious how the spores are discharged successively, as Buller describes. One would expect the contents of the basidium to escape through the hole in the sterigma as soon as the first spore was shot off; either the hole is too minute or it contracts elastically.

The small basidia and their appendages develop in a similar way; their contents are never coarsely granular or guttulate. The intermediate basidia, which are scarce, develop at intermediate depths in the subhymenium and bear spores of intermediate size. I have never seen small and large spores on the same basidium.

The motor mechanism of the fruit body

One must distinguish between the chief motor mechanism which expands the fruit body and the expansion of the palisades which merely sets up local tissue pressures. The chief motor mechanism lies in the medullary hyphae which gradually inflate acropetally in a grand period of growth in the stem, pileus, and trama, as I have described in Collybia apalosarca. Inflation in the primordium is inconsiderable until the pileus is initiated: the medullary hyphae of the stem then rapidly inflate carrying up the rudimentary pileus, the central medullary tissue of which also begins slowly to inflate. The wave of inflation passes from the stem into the pileus, radially to its margin and obliquely outward and downward in the trama of the gills: the stem is nearly fully expanded when the pileus is about halfexpanded. The cells in the primordial shaft average $20-2.5 \mu$ on delimitation; they straightway begin to inflate slowly and in the mature stem they average $400 \times 20 \mu$, elongating about twenty times; this proportion corresponds with the macroscopic growth of the stem: the primordial shaft reaches an average height of 2 mm. and the stem of the full-grown fruit body averages 5 cm.

Not all the medullary hyphae inflate. Very few inflate in the core of the primordial shaft, which is thus disrupted to form the hollow of the stem. The inflating hyphae are dispersed evenly through the medulla of the pileus and trama, which do not become hollow.

The inflation of the palisade hyphae does not take place in such uniform sequence. It is generally basipetal, beginning in the apical cell, soon after it has dropped behind the growing margin, and extending through two to four subterminal cells. The terminal cells of the scattered cortical filaments on the stem, the cells of the pile on the pileus, and the basidia all begin to inflate before their corresponding medullary hyphae, and are often nearly fully inflated before any appreciable internal enlargement has occurred.

Whereas the development of the stem is really indirect (a period of apical growth as a primordial shaft being followed by a distinct period of inflation), the limb and gills develop directly, inflation of the cells following closely on delimitation. The limb is at first convex with slightly incurved margin because, as already explained, the hyphae of the pile inflate more rapidly and to a greater extent than the hymenial hyphae. The limb then straightens centrifugally through the inflation of its medullary hyphae and, finally, owing to the intercalary growth of the hymenium and inflation of the gills, it may be thrust upward, becoming concave and the pileus more or less infundibuliform. The pileus mechanism is similar to, but the inverse of, the apothecial mechanism(3).

THE FRUIT BODY OF HYGROPHORUS HYPOHAEMACTUS

Apart from slight differences in the spores, basidia and cystidia, the fruit body of *H. hypohaemactus* is very like that of *H. firmus*: it is gymnocarpic with exogenous pileus, has both inflated and uninflated medullary hyphae, a distinct pile at least over the centre of the pileus, no pile or palisade on the stem, a sterile gill edge, and the pigment in the cell sap. It differs as follows:

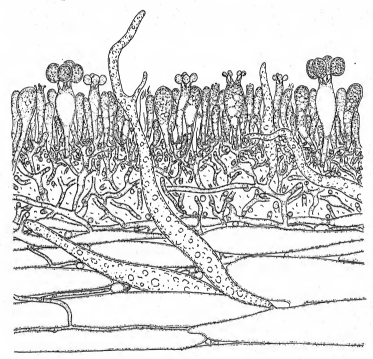


Fig. 5. Part of a longitudinal section of the gill of H. hypohaemactus, showing the "pseudocystidia"; \times 500.

(1) The hyphae of the pile do not inflate, being $2-7 \mu$ wide in mature fruit bodies, but their walls become very mucilaginous. They are sparingly branched, clamped and with simple ends, and are more or less perpendicular to the surface over the disc and decumbent over the limb. The thick grey mucilage, derived from their walls, becomes so copious in wet weather that it swells beyond the limits of the pileus in dentate and appendicular processes.

(2) The superficial hyphae of the stem also become mucilaginous

and are not inflated (2-5 μ wide).

(3) The subhymenial hyphae have thin mucilaginous walls and

are not inflated (2–5 μ wide), so that the subhymenium is subgelatinous and rather thick (ca. 50 μ at the base of the gills, 10–20 μ

near the edge of the gills, and ca. 40 μ on the sides).

(4) Some of the laterals of the tramal hyphae do not contribute to the hymenium with clusters of basidia but lie obliquely disposed as elongate fusiform cells with densely granular oleaginous contents and simple, rarely furcate, obtuse ends. They are either wholly embedded, passing from the trama to the subhymenium, or they project up to 50 μ beyond the hymenium into the gill space or from the edge. They are straight or sinuous, simple or once or twice branched, aseptate, thin-walled, $50-600\times4-16$ μ , $\times3-6$ μ at the apex, and they look like tramal cystidia. The shorter ones may arise from the subhymenial hyphae and appear as true cystidia in the hymenium. In some fruit bodies they are much more abundant than in others, and owing to their varied origin and disposition they can hardly be called pleurocystidia.

The size of the fruit body

The fruit body varies greatly in size in the varieties of H. firmus, from 5 mm. to 17 cm. high, and, as I have shown in connection with Collybia apalosarca, the problem is to determine whether this variation is due to differences in the amount of inflation or of apical growth of the hyphae. As in C. apalosarca, I have analysed a sufficient number of fruit bodies to show, I think, that it is mainly due to differences in apical growth, since the average amount of inflation is roughly the same. Error is, of course, possible in this method of averaging and extrapolating. Only a few cells are measured in the middle section of the stem, midway between apex and base, and these are assumed to indicate the degree of inflation of the whole fruit body. Fortunately the number of gill ranks gives an independent and simple check on the state of the pileus; this indicates that the pilei of those varieties with small fruit bodies are juvenescent. Also, in the same fruit body, the cells vary greatly in their degree of inflation, and owing to the practical difficulty of tracing the longer cells, the shorter are liable to be picked for measurement. To avoid choosing, I seize upon the septa, measuring the cell on each side of it and then pass to the next obvious septum and so on, finally taking the average of fifty. Nevertheless, the results from both Hygrophorus firmus and H. hypohaemactus are consistent enough to prove that one is not far from the truth. The fruit bodies of varieties sericeus, minimus and gracillimus may be taken as juvenescent compared with the typical fruit body of H. firmus var. militaris, and likewise those of H. hypohaemactus, but those of the varieties longipes and pachyphyllus are clearly overgrowths (the average length of an inflated cell from the stem is 400 μ). But the variations in size between fruit bodies of the same

troop, i.e. contemporary from the same mycelium, are as often due to differences in apical growth as inflation.

Table A. Showing the relation between the size of stem and inflation of the cells in H. firmus

		n size mm. Width in middle	Stem- length ratios	Average cell length from middle section of stem	Ratios of average cell lengths	Cell extremes	
H. firmus var. militaris	53 50	2·5 2·5	1 0•94	460 429	0.0	92-1513 150-825	One tuft
H. firmus var. stratiotes	86 75	4 5	1·6 1·4	563 584	1·3	170-1170 200-1750	One tuft
H. firmus var. gracillimus	17 15	I I	0.3	398 374	o·9	150-750 120-720	One tuft
	11 12	I	0.5	325	0·7	100–850 120–600	
H. firmus var. stenophyllus	40 30	5 4·5	o·75 o·6	453 417	0.0	150-900 150-750	One tuft
H. firmus var. longipes	130 155	5 7	2·5 2·9	407 470	0.9	150-800 120-1340	One tuft
H. hypohaemactus	37 35 25 20	3 3 2·5 2	0·7 0·5 0·4	430 515 435 400	0.8 1.1 0.8	100–1000 100–1300 100–1050 175–850	One tuft

The gill arrangement

Since the gills subsequent to the primaries are regularly intercalated at the margin of the limb as the pre-existing gills diverge and before the inflation of the medulla, the number of gill ranks roughly measures the extent of marginal growth; other things being equal, the greater the extent of marginal growth the more the gill ranks, while the amount of inflation is secondary and immaterial to the origin of the gills. Analysis of the gill arrangement provides a convenient check on the direct analysis of the size of the fruit body. The data obtained from about eighty fruit bodies are enumerated in Table B.

The table shows two facts. Firstly, in those varieties with small fruit bodies, namely sericeus, minimus and gracillimus, the small size of the pileus is due to the lesser growth of the limb. This is the explanation deduced in the previous section for the shortness of the stem in var. gracillimus, and one is therefore justified in considering the fruit bodies of these varieties as juvenescent. Secondly, the variation in size of the fruit bodies growing in the same tuft is due partly to a lesser extent of apical growth and partly to the lesser inflation. When many primordia develop close together from the same my-

Table B. Showing the relation between the gill arrangement and size of fruit body in H. firmus

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celium, they grow equally for a short time, then some take the lead, developing normally, and the rest are more or less stunted: in fact, in caespitose species one can generally find in the mature tufts all transitions from fully developed fruit bodies to rudimentary and distorted primordia, and care must be taken not to include these abortive fruit bodies in material for developmental studies. Sometimes the stunted primordia have undergone their full amount of apical growth, for they possess the full complement of primaries and

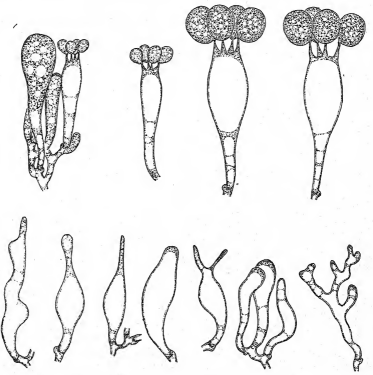


Fig. 6. Large and small basidia and cheilocystidia of H. hypohaemactus; × 1000.

gill ranks, but they are stunted through lack of water for inflation. Apical growth of other primordia may be checked prematurely and these develop into fruit bodies which are not merely small but which have fewer primaries and gill ranks. The mycelium lays down more initials than it can mature; their demands exceed the supply of food and water, and but a few develop normally. It is not until one begins to watch the primordia in the field that one realises how many failures there are. I have never seen an aborted primordial shaft expanded into a *Clavaria*-like body; unless the pileus is developed to an appreciable extent, the primordium aborts completely.

Discussion on development

Some American investigators have contended that the gills of agarics arise through the buckling of the hymenium as it undergoes intercalary growth and not by an active outgrowth along certain paths(2). They state that the hymenium of the primordium is at first flat and, as the basidia inflate and as more are intercalated sympodially, so the hymenium is compressed between the incurved margin of the limb and the stem apex: a tension develops which is relieved by throwing the hymenium into radial folds. That this explanation rests on misunderstanding is evident from the following considerations:

(1) It is not obvious that such a state of tension would be relieved by simple radial buckling. Considering the intricate connections of the hymenium it would most probably be thrown into anastomosing wrinkles, defining more or less hexagonal areas, e.g. the species of Cantharellus, Craterellus, Cyphella, Marasmius, Campanella and Stereum

with wrinkled hymenium.

(2) Why should not such a lateral pressure cause merely a hyponastic curvature of the limb, which is free to move in either direction? Actually the tension in the primordial hymenium, when the gills are developing, is never enough to overcome that of the pile on the pileus which forces the limb towards the stem.

(3) If the gill ridges buckle out from the limb, why are there not schizogenous cavities overlaying the gills, like the hollow in the stem?

(4) Why are gill ridges not developed on the upper side of the pileus in species with a compact pile or palisade, e.g. species of Boletus, Pluteus, Mycena, Marasmius, Psathyra, Coprinus, etc.? Such a palisade of contiguous inflated cells, increasing their numbers by sympodial branching, presents exactly the same problem as the primordial hymenium. At most, anastomosing wrinkles develop, as in Pluteus phlebophorus, Lactarius fuliginosus and Psathyra corrugis.

(5) How can the secondaries and higher ranks be intercalated regularly by this means? When as rarely happens anastomosing

wrinkles are developed, they offer no such order.

(6) How would the tramal hyphae stand the stretching? For the cells of some might be pulled out for several millimetres, and if they broke that part of the hymenium supplied by them would perish.

(7) Why should the tramal hyphae be so compactly arranged and

strictly longitudinal in the typical gill?

I think that it will always be found that the gill ridges are lines of outgrowth as I have described in *Collybia apalosarca* and *Hygrophorus firmus*. But how they are blocked out at the margin of the limb by sectors of inhibition is a profound problem. The morphologist always comes against such barriers which limit his researches and show where he must hand over to the experimenter.

The biology of the fruit body

The fruiting of Hygrophorus firmus in Malaya is seasonal. It takes place during a week or two in each wet season after a dry spell, which must be of two or three weeks duration and pronounced enough to dry off the surface layer of humus in the forest. In Singapore such a spell generally occurs twice a year, in February and in July or August, and it is followed by a period of three to four months' intermittent rain during which the humus is always moist. The fruit bodies of H. firmus then begin to develop some six weeks after the break in the dry weather. Over a small area of forest where the rainfall is uniform, one may therefore find the troops of fruit bodies twice a year during one or two weeks, and only very sporadically, if at all, in the intervals. Each mycelium fruits once only in each season.

I have watched to completion the growth of forty-three fruit bodies of *H. firmus stratiotes*, and many others which were less complete.

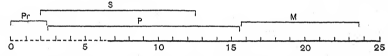


Fig. 7. A diagram of the life of a fruit body of *H. firmus* var. stratiotes: the continuous basal line covers the period of sporing: the figures are days (24 hours): *Pr*, the primordial shaft; *S*, stem elongation; *P*, pileus-enlargement; *M*, maturity.

Development is gradual. The total life of the fruit body varies from twenty to twenty-seven days, being generally twenty-three to twentyfour days. The apical growth of the primordial shaft occupies the first two or three days. The stem elongates during the next seven to twelve days; the pileus continues to expand for a further three to five days and the fruit bodies persist full-sized and mature for a final period of four to ten days. Sporing begins when the pileus is 4-6 mm. wide, that is to say on the sixth to seventh day. These facts are delineated for an average fruit body in Fig. 7. It is not, of course, easy to judge either when the primordium is initiated, because it is hidden in the soil, or when the fruit body is thoroughly effete, because the limb collapses gradually: a day or so may therefore be added or subtracted both from the beginning and the end of this scheme. But, certainly, the primordial shaft grows remarkably quickly. The stem elongates most rapidly during the interval from the seventh to tenth day, the pileus from the ninth to thirteenth day. The stem often begins to rot at the base before the limb collapses, especially if there is much rain, and this may cause the premature death of the fruit

The fruiting of H. hypohaemactus also appears to be seasonal, but

owing to its scarcity I cannot state this categorically. During the four years, 1929–32, it has fruited twice a year in the Singapore Botanic Gardens at intervals of two to four months after the break of the dry spell. It certainly develops much later in the wet season than H. firmus. General observations show clearly that the fruit bodies have a similar slow growth and long period of maturity, although I have not been able to study them in detail. This slow development and indefinite period of sporing is probably characteristic of the more massive gymnocarpic agaries such as Hygrophorus, Clitocybe, Entoloma, etc., and it is in striking contrast with the rapid evolution and precise mechanism of Collybia apalosarca and the angiocarpic genera Amanita or Coprinus.

Systematic section

Hygrophorus firmus B. & Br. (=H. firmus var. typicus).

Pileus 1-4 cm., at first convex, becoming plane and slightly umbilicate, sometimes infundibuliform or pervious to the base of the stem, minutely scurfy squamulose over the disc, innately fibrillose and striate toward the margin, dry, orange-red or scarlet, paler on expansion and pale yellowish when old; margin slightly incurved at first, often crenatoplicate.

Stem 3-8 cm. × 2-12 mm., equal, slightly thickened above or attenuate downward, dry, smooth, becoming hollow, rather stiff, often flattened below, paler con-

colorous, discolouring pale yellowish; base white, abrupt.

Gills shortly decurrent, often more or less broadly sinuate, distant, 1-4 ranks, with 16-40 primaries, 2-3.5 mm. broad, rarely forked near the stem, rather thick, not veined or very slightly, concolorous at first, becoming pale yellow or whitish, yellowish at the base; trama pallid yellowish.

Flesh thin, 1-2 mm. in the centre, 0.3-0.5 mm. half-way to the margin, watery,

rather soft, concolorous; smell and taste none.

Spores white, ellipsoid, smooth, thin-walled, dimorphous: large spores $12-16 \times 7-10 \mu$, broadly oblong ellipsoid, ends blunt, slightly flattened adaxially, sometimes narrowed distally rather suddenly and waisted, with a prominent lateral-basal apiculus, with dense granular-oleaginous contents: small spores $5-8 \times 3\cdot 5-4\cdot 5 \mu$, more or less pip-shaped, widest in the proximal half, with hyaline, cloudy vacuolate contents.

Basidia dimorphous: large basidia 50-75 \times 12-16 μ , clavate, with dense granularoleaginous contents at first, becoming hyaline, with four large sterigmata 8-11 \times 3-4 μ at the base: small basidia 28-40 \times 6-8 μ , subclavate or subcylindric, contents cloudy-vacuolate, with four, rarely two, sterigmata 5-7 \times 1.5-2 μ at the base.

Cheilocystidia $25-60 \times 7-30 \mu$, cylindric, clavate or vesicular, rarely ventricose, thin-walled, colourless, vacuolate, sometimes with one to five short cylindric processes up to 5μ long from the distal end, as abortive sterigmata: forming a narrow sterile edge, often inconspicuous and with all transitions to sterile basidia.

Caulocystidia absent or as a few irregular sterile basidia at the apex of the stem. Habitat, in troops or small tufts in the lowland and mountain forest up to

4000 ft.; Ceylon, Malaya.

The varieties of *H. firmus* can be arranged in two sections, Ovalisporae and Macrosporae. I have given, in the following descriptions, only the distinctive characters of the varieties which in other respects, unmentioned, may be taken as identical with the typical state.

Ovalisporae. Sporis magnis $12-16 \times 7-10 \mu$, late ellipsoideis vel ovoideis, apicibus obtusis, vix attenuatis.

var. militaris var.nov.

Stipite albo vel primo pallide flavo.

This variety appears commoner than the type, at least in Singapore. Growing in troops, with scarlet cap and stiff white stem, its carriage is soldierly.

var. puniceoides var.nov.

Pileo 7–8 cm. lato, magno, convexo-plano, non umbilicato: stipite 6–7·5 cm. \times 7–9 mm., claro flavo: lamellis latis, 6–10 mm., adnexis, saepe dentibus parvis decurrentibus praeditis, pallide citrino-flavidis: carne crassa, 3–4 mm. medio pileo, pallide alba: sporis magnis minusculis, oblongis, saepe paulo constrictis, 12–15 \times 7–9 μ .

I found this variety once, at Tembeling in Pahang, November 4th, 1930. Macroscopically it is so like *H. puniceus* that one would not consider it related to *H. firmus*, although microscopically it is essentially the same.

Macrosporae. Sporis magnis $16-25\times7-11~\mu$, longo-ellipsoideis, plus minus fusiformis, saepe paulo curvatis vel allantoideis, apicibus subacutis, distincte attenuatis.

var. stratiotes var.nov.

Forma typica persimilis sed sporis majoribus, $18-25 \times 7-11 \mu$.

Macroscopically this variety is indistinguishable from the typical state. The basidia are generally of the same size, though the large ones may reach 18 μ wide. It is the commonest variety in Malaya, and I have found it frequently in Singapore, Johore, Pahang and Negri Sembilan, where it may be expected every season.

var. pallidipes var.nov.

Stipite albo.

This variety stands to var. stratiotes as var. militaris to the typical state. It is not uncommon.

var. depallens var.nov.

Pileo albido vel flavido, raro carneo-flavido: stipite 7-10 cm. × 4-6 mm., flavido, apice albo, vel ex integro albo: lamellis latis, 6-7 mm., pallide carneo-ochraceis.

I have found this variety only in the Reservoir Jungle in Singapore. It is poorly pigmented and the pinkish gills suggest *Entoloma* or *Leptonia*.

var. sericeus var.nov.

Pileo parvo, 5–12 mm., cano-albo, subfibrilloso, sicco atomato: stipite 2–2·5 cm. \times 1·5–2 mm., ceraceo, citrino-flavido: lamellis albis dein pallide carneis: sporis magnis 16–21 \times 6·5–9 μ : sporis parvis 6–8·5 \times 3–3·5 μ .

I have found this variety on the hill, Bukit Timah, in Singapore. Macroscopically it exactly recalls *Leptonia sericella*: the loose, sparingly branched, decumbent hyphae give the limb the silky fibrillose appearance. It differs from var. *depallens* in the smaller size, narrow gills, smaller spores and paler colour.

var. flavus var.nov.

Stipite et pileo citrino-flavo vel saturatiori: lamellis flavidis, dein albidis: sporis magnis 16-22 × 10-11 μ : sporis parvis 5-7 × 3·5-4·5 μ .

This variety appears every year in the Aroid Rockery in the Singapore Botanic Gardens, where there are many forest trees. The fruit body is brilliant yellow without a trace of red.

var. flavo-albus var.nov.

Pileo flavido: stipite albo: lamellis albis, latis, 4-6 mm.

This variety is so very close to the preceding that, perhaps, it hardly deserves a name. It is a further step, however, to the following albino condition. I have found it in the Reservoir Jungle, Singapore. Bresadola's illustration of *H. lucorum* (*Iconogr. Mycol.* vII, 314) would pass macroscopically for this variety.

var. albus var.nov.

Ex integro albus.

I found a troop of fourteen fruit bodies of this variety by the Cheka River in Pahang, on November 12th, 1930. The fruit body is devoid of pigment, though in shape, size, and microscopic characters identical with var. stratiotes.

var. roseus var.nov.

Pileo purpureo-roseo vel carneo: stipite albo vel pallide carneo: lamellis pallide carneis, pallescentibus.

This variety is not uncommon. I have found it in Singapore and at Tembeling and Fraser's Hill (4000 ft.) in Pahang. The fruit bodies are often rather larger than typical, especially in length of stem. It leads both to the following and to var. longipes.

var. amethystinus var.nov.

Pileo purpureo-vinaceo: stipite albido: lamellis claro amethystinis, ut Laccaria laccata var. amethystina.

I found this most striking variety as a troop of twenty fruit bodies in the forest at Tembeling, Pahang, on November 9th, 1930. The

colour is so unlike that of a *Hygrophorus*, lacking all trace of red and yellow, that it suggested *Tricholoma* or *Clitocybe*. Microscopically it is indistinguishable from var. *stratiotes*.

var. pachyphyllus var.nov.

Pileo 5-6 cm. lato: lamellis perlatis, crassis, 8-10.5 mm. latis, ad basim 2 mm. crassis.

I have not analysed the property of thick gills in this variety, but it is doubtless caused both by an excessive marginal growth and inflation or intercalary growth. It occurs not infrequently in the Reservoir Jungle, Singapore.

var. stenophyllus var.nov.

Pileo convexo-plano, citrino-flavo: stipite citrino-flavo: lamellis albis, angustis, 2-2·5 mm. latis, subdecurrentibus, late sinuatis vel subarcuatis.

I found a troop of some fifty fruit bodies of this variety on Fraser's Hill, Pahang, May 27th, 1930, at 4000 ft. altitude. Only the largest fruit bodies had spores; those with pilei less than 16–20 mm. wide were not yet fertile. It may be an abnormal form of var. flavus, partly sterile and thus with narrow gills.

var. longipes var.nov.

Stipite saepe altissimo, 5-17 cm., apice 4-7 mm., basi 5-10 mm., albo: lamellis saepe perdecurrentibus, 3·5-4·5 mm. latis, albis.

The pileus is typical, 2.5-4.5 cm. wide, scarlet fading to orange or pale yellow, and with the full gill complement. The enormous length of stem, as shown in the preceding section, must be due to long-continued apical growth of the primordial shaft, since the average inflation of the cells is typical. Such a state might be evoked through an external factor as weak light, unusual temperature or high humidity, but the fruit bodies may be found in open sunny places by paths in the forest, in deep shade, in humus, or on bare sandy banks, and both in the lowlands and the mountains. I have found it at Padang Piol, near Tembeling, and at Fraser's Hill (4000 ft.) in Pahang. In the collection from Padang Piol and most of those from Fraser's Hill the spores are as in var. stratiotes, but in one collection from Fraser's Hill they were unusually large, as shown in Fig. 8 e: the large spores measured $22-27 \times 9-11 \mu$ and the small spores $9^{-12} \times 4^{-4.5} \mu$. Scattered through the forest on Fraser's Hill I have also seen specimens of this variety which were wholly white or with the faintest tinge of yellow on the pileus, but I have no accurate data on them.

var. minimus var.nov.

Pileo pusillo, 4-10 mm. lato, convexo-plano, raro subumbonato vel subum-

bilicato: stipite brevissimo, $5-10\times1$ mm., solido: lamellis arcuatis, perdecurrentibus, distantis, 7-10 primis, ordine uno, raro secundo incompleto, instructis.

Apart from their minute size, the fruit bodies of this variety are identical with those of var. *stratiotes*. As shown already, they must be regarded as juvenile forms with limited apical growth and normal turgescence. It is not infrequent, but easily overlooked. I have found it in Singapore, round Gunong Panti in Johore, and at Tembeling in Pahang. Generally the fruit bodies are scattered and few.

var. gracillimus var.nov.

Pileo pusillo, 4–7 mm. lato, convexo-plano, subumbonato, plus minus turbinato, flavido vel albido, centro saturatiori carneo-aurantiaco: stipite gracili, 15–25 × 1 mm., solido, translucido, pallide aurantiaco: lamellis subtriangulis, breviter decurrentibus, uno ordine, raro secundo incompleto, instructis, 10–12 primis, 1–1·5 mm. latis: sporis 18–23 × 9–10, et 8–10 × 3·5–4 µ.

This variety differs from the preceding in the longer, slender stem and pale colour. It looks like an *Omphalia*. I have found it only on Fraser's Hill by the path to the waterfall, but in a troop of over a hundred individuals growing in moss on a sandy bank, and in two successive seasons, in May and November, 1930.

Hygrophorus hypohaemactus spec.nov.

Pileus 2-2·7 cm. latus, convexus dein planus, margine primo incurvato, viscoso-papillosus, glutine hyalino griseo viscido adnato, in medio pileo 1-1·5 mm. crasso, marginem versus tenuiori, saepe supra marginem in vittas triangulas usque 1 mm. longas projicienti, obtectus, umido striatus, sub glutine rubro-sanguineus, dein pallescens, demum pallide aurantiacus vel flavidus.

Stipes 3-6 cm. × 2·5-3·5 mm., cylindricus, solidus dein cavus, viscidus, glutine simili, saepe peronato-disrupto vel crispato, obtectus, pallide aurantiacus vel rubro-

aurantiacus, demum pallescens.

Lamellae adnatae vel adnexae, separantes, subdistantes, crassiusculae, ordinibus 3-4 instructae, primae 18-20, 2·5-3·5 mm. latae, ceraceo-submucosae, margine obtuso continuo dein mucoso et interrupto praeditae, albae, basim versus pallide aurantiacae.

Caro tenuis, medio pileo 1-1.5 mm. crasso (glutine subtracto), concolor: odore

deficien

Sporae albae, laeves, dimorphae: sporae magnae $8-10\times6.5-8.5$ μ ovoideae vel lato-ellipsoideae, intus granuloso-oleaginosae: sporae parvae $4-5\times3-3.5$ μ , sub-globosae, basim versus attenuatae, intus nebuloso-vacuolatae, vix vel haud granulosae.

Basidia dimorpha: b. magna 33-44 × 10-12 μ , stipitato-ventricosa sterigmatis 4, 5 μ longis, praedita: b. parva 20-28 × 5-6 μ , subclavata, sterigmatis 4, 3-3-5 μ

longis, praedita.

Cheilocystidia $16-40 \times 4-9 \mu$, polymorpha, subclavata, clavata, ventricosa, saepe flexuosa, hyalina, vacuolata, tenuiter tunicata, apicibus plerumque simplicibus, saepe 1-ramosis vel lobatis vel appendicibus 2 sterigmatoideis praeditis: pleurocystidia vera absentia.

Hab. ad terram in silvis: Malaya.

I have found this species only in the Garden's jungle and once at Tembeling, in Pahang. It has caulocystidia only at the extreme apex of the stem where they are like the cheilocystidia but rather longer, up to 60μ , more flexuous and more branched and irregular, and they form a few compact clusters, as in *H. firmus*, immediately preceding the hymenium.

Systematic discussion

It is not easy to decide in a species so variable as Hygrophorus firmus which state shall be reckoned the typical. Very fortunately the collections from Ceylon, whence the species has hitherto been known, are the most generalised and approach nearest to what is supposedly the ancestral state. They are thus qualified on grounds other than mere "type collection" to hold such a position. The same state occurs in Malaya, and with the local material I have amplified Petch's description in the foregoing section as H. firmus var. typicus. These dimorphous species are surely derived from the section of Hygrophorus including H. miniatus, H. Reai, H. turundus, etc., which have fruit bodies of similar shape, colour and manner of development. The spores of this section are typically subglobose, $6-9 \times 4-7 \mu$, with fairly dense, granular or cloudy vitreous contents, and not only do the large spores of H. firmus pass through such a stage in their development but those of *H. hypohaemactus* are little removed from it and lead through the Ovalisporae to the Macrosporae (vide Fig. 8). The fruit bodies of this section are also of medium size and more or less fully pigmented as in H. firmus var. typicus. Thus the other states of H. firmus may be looked upon simply as variants in size and pigmentation. Regarding these varieties there are the following facts:

(1) In any one troop the fruit bodies, which are presumably derived from the same mycelium, all have the same special characters. I have never seen two varieties mixed in the same troop or even

growing in the immediate vicinity of each other.

(2) The same varieties occur in such widely separate localities as Singapore, Johore and Pahang, e.g. stratiotes, militaris, minimus, palli-

dipes, roseus and longipes.

(3) Certain varieties I have found repeatedly on the same patch of ground each season so that their mycelia must have constant characters, e.g. militaris, stratiotes and flavus in the Singapore Botanic Gardens (1929–32), pallidipes in the Reservoir Jungle, Singapore (1930–32), and gracillimus and longipes on Fraser's Hill, Pahang (May,

November, 1930).

The characters of the varieties are clearly fixed and hereditary, and it is a problem whether some should not rank as species. Indeed, should not one propose a new genus based on the dimorphous spores, with two subgenera, the one for *H. firmus* with dry pileus and stem and the other for *H. hypohaemactus* with viscid? Yet it is equally clear that many of the varieties are merely colour forms and that intermediates will probably be found. Moreover, anatomical knowledge

of the Hygrophorei is far too meagre to allow the recognition of natural genera. Is it usual in *Hygrophorus* for some only of the hyphae to inflate the fruit body? I would have refrained even from varietal names were they not necessary for accurate description and to emphasise the extraordinary range in size, shape and colour which *H. firmus* displays. And though it is a common species, many years must elapse before we have a full knowledge of it: the period of fruiting is so brief that one traverses the forest without meeting with it many times. The following remarks therefore are only tentative.

The Ovalisporae may differ specifically from the Macrosporae. The spores of *H. firmus* are fusiform-ellipsoid because the stiffening of the distal part of the wall, when the spore is in the subglobose

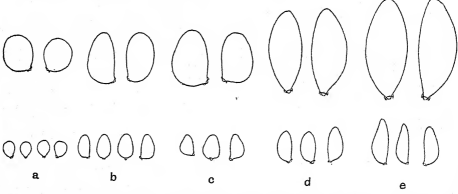


Fig. 8. Large and small spores: a, H. hypohaemactus; b, H. firmus var. puniceoides; c, H. firmus var. typicus; d, H. firmus var. stratiotes; e, H. firmus var. longipes (Fraser's Hill, vide text); × 1000.

stage, is delayed and this part is further protruded into a subacute apex through the pressure in the basidium, and the process is carried farthest in the Macrosporae. In the Ovalisporae, var. puniceoides might be separated specifically from H. firmus var. typicus and var. militaris on the shape and colour of the fruit body and the rather smaller spores, which are most like those of H. hypohaemactus. In the Macrosporae, probably only var. gracillimus deserves specific rank. The varieties longipes, minimus and pachyphyllus are only growth forms. The remainder are chiefly colour forms in which the pigment is either limited to certain parts or modified into yellow, pink or purple. The varieties stenophyllus and sericeus combine differences both in colour and form.

H. similis Petch is macroscopically very like H. firmus, but its spores are monomorphous and narrowly ellipsoid or subcylindric, $6-9 \times 3-3 \cdot 5$ (-4) μ . I have examined part of the type collection, Herb.

Perad. 5580. It seems very close to H. Reai, but I could make out no

structural details in the dried specimen.

Finally it must be remarked that Petch's statement that the large spores of *H. firmus* are verrucose is an error. I have examined Herb. Perad. 2299, determined by Petch, and find the spores smooth. It is well known that a densely granular cytoplasm may give the appearance of a rough epispore when the spore wall is very thin and transparent, and an immersion lens is needed to decide the point.

Summary

Hygrophorus firmus and H. hypohaemactus sp.nov. form two kinds of basidiospore. In the same fruit body they produce large spores with dense contents on large basidia, and small spores of half the linear measure and with vacuolate contents on small basidia. The meaning of the dimorphism is unknown.

Both species occur in Malaya, H. firmus being known previously from Ceylon. H. firmus in Malaya is exceedingly variable. Sixteen varieties are proposed under two sections according to the spore character: most are colour varieties but some are shown to be juvenile forms, others to be overgrowths. H. hypohaemactus is rare.

H. hypohaemactus differs from H. firmus in the viscid pileus and stem and in the smaller less dimorphous spores. Both are related to the group of H. miniatus, H. turundus, etc. The systematic position is discussed.

The structure, development and variation in size of the fruit body

are explained in detail.

The fruit body of *H. firmus* is gymnocarpic with exogenous pileus. There is a pile on the pileus, at least over the disc, but not on the stem. The gill edge is sterile with cheilocystidia: there are no pleurocystidia. Development is gradual, occupying about a fortnight. The growth of the primordial shaft takes two to three days. The stem and pileus slowly inflate acropetally during the next ten to twelve days. The mature fruit body lasts about a week, making the total life a little more than three weeks. Sporing begins about the sixth day when the pileus is 4-6 mm. wide. These observations were made in Singapore at a mean temperature of 80° F. (70-90° F.).

The limb and gills develop by apical growth of the hyphae followed by acropetal inflation, exactly as the primordial shaft. In the stem and pileus many hyphae do not inflate but are passively drawn out. Both kinds of basidium are borne on the same hyphae, the large basidia being the first formed and arising deeply in the subhymenium. The large spores are carried beyond the level of the

small spores. The hymenium is aequihymeniiferous.

The fruit body of H. hypohaemactus is essentially similar, but the

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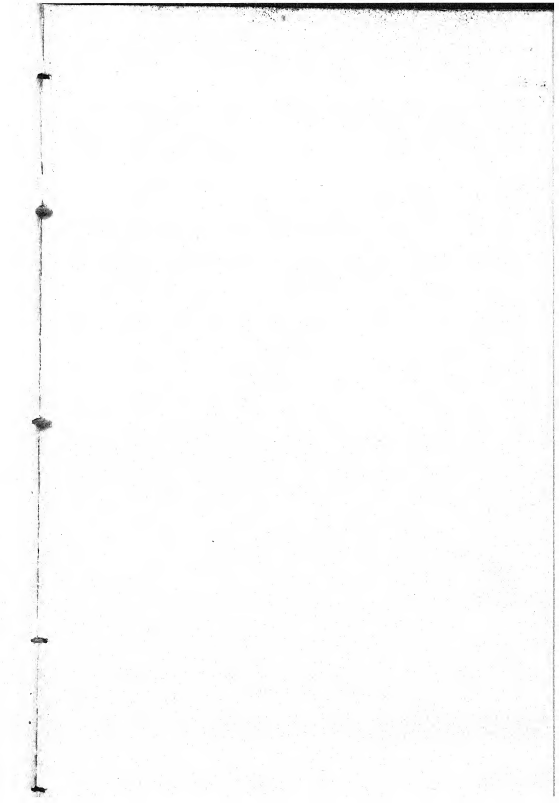
hyphae of the pile on the pileus and those on the surface of the stem and in the subhymenium have mucilaginous walls, and there are peculiar hypha-like tramal cystidia with oleaginous contents in the gills.

I must, in conclusion, express my obligation to the Director of Agriculture, Ceylon, for the loan of authentic material and drawings of the several species of Hygrophorus.

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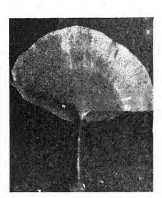


Fig. 1.



Fig. 2.

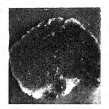


Fig. 3.



POLYSTICTUS XANTHOPUS

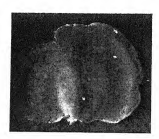


Fig. 5.



Fig. 6.

POLYSTICTUS FLABELLIFORMIS

A NOTE ON THE VARIATION OF PORES OF POLYSTICTUS XANTHOPUS FRIES AND POLYSTICTUS FLABELLIFORMIS KL. AT HIGH ALTITUDES

By S. R. BOSE

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(With Plate II)

The hymenial surface of specimens of *Polystictus xanthopus* Fr. at high altitudes shows three different kinds of porous areas, viz. some with typical very small pores, some with much bigger pores and some with hydnoid pores (Pl. II, figs. 1, 2 and 3). In one piece of dead branch of a tree, specimens with typical pores and hydnoid pores were growing not very far off from each other (Pl. II, fig. 4), and so it is concluded that they are variations of one and the same species. Such variations are hardly found in the plains where the fungus is abundant, with very small typical pores on the hymenial surface.

Specimens of *P. flabelliformis* KI. similarly collected from high altitudes show two kinds of porous surface, some with very small pores and others with much bigger pores (Pl. II, figs. 5 and 6). In other respects, morphological and anatomical, the specimens are exactly similar. In the plains we always get them with small pores

on the hymenial surface.

This illustrates, as has been noted by French mycologists (Sauger, Josserand, Maire, etc.), that the dimension of the pores in the hymenial surface—one particular character—is not always a safe

guide in the delimitation of species.

The specimens described in this note were collected from high hills of four different localities: (1) Cherrapunji, Assam, by me (8000 ft. elevation) in March, 1929; (2) Lokra hills (8000–10,000 ft. elevation) by Dr N. L. Bor in January, 1934; (3) Bhutan border (10,000 ft. elevation) by Mr K. P. Biswas, Curator of the Shibpur Herbarium Royal Botanic Gardens, in April 1934; (4) Pareshnath hills (2000 ft.) by Mr K. P. Biswas in November 1934.

NOTE ON ABNORMAL SPORES IN PODOSPORA MINUTA

By WINIFRED M. PAGE, M.Sc.

(With 5 Text-figures)

 $P_{ extit{ODOSPORA MINUTA}}$ occurs commonly on rabbit pellets. It appears early in the succession of fungi, usually fruiting at the same time as Ascophanus carneus and other Discomycetes.

The spores germinate readily and single-spore cultures are easily

obtained.

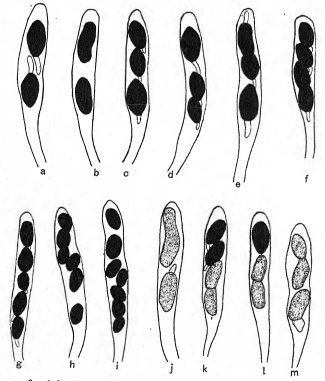


Fig. 1. Types of asci showing variation in number of spores and abnormal spores. $\times 200$.

The normal asci contain four spores, each of which is capable of producing a mycelium giving rise to fruits (Fig. 1f). In perithecia grown in culture and also in those growing on rabbit pellets collected

from various sources, asci with abnormal spores sometimes occur. From one to seven spores which, apart from size, are normal in appearance have been observed (Fig. 1 a-i). In addition irregular spores, some of which never completely darken, are occasionally

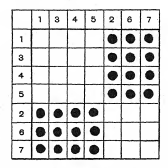


Fig. 2. Table showing results of crossing mycelia from A and B spores.

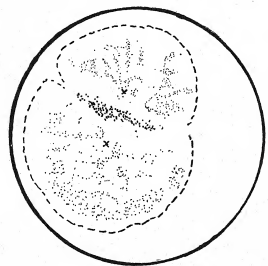


Fig. 3. Diagram from photograph of culture from A and B spores. Points of inoculation indicated by ×. The fine dots show abortive perithecia and the broken lines the limits of the mycelia.

seen (Fig. 1j-m). Of asci with other than four spores, the one giant and two normal is the most common variety (Fig. 1c-e). Asci with two dwarf and three normal spores are not so frequent, but a number of the dwarf spores have been isolated and germinated (Fig. 1g). As in Sordaria fimicola (four-spored form) the perithecium initials in

the mycelia from these spores never mature. When the spherical stage is reached no further development takes place (4, 5, 6).

The mycelia from these dwarf spores were found to be of two different, but complementary, strains. A number of crosses were

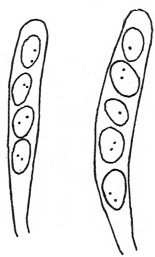


Fig. 4. Sections of young four- and five-spored asci showing nuclei. × 400.

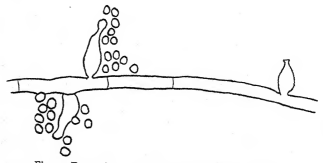


Fig. 5. Formation of small colourless spores. × 1600.

made and the results are shown in the table (Fig. 2). The details of the results agree with those of Sordaria fimicola (6).

The normal spores contain two nuclei and the dwarf spores one

nucleus each (Fig. 4).

The eight-spored form of this fungus is not common; I have found it only once in many years of collecting. It was, however, cultured. Each of the eight spores is uninucleate and capable of producing mycelia which fruit.

In the four-spored variety very tiny colourless spores are produced, being cut off from flask-shaped outgrowths (Fig. 5). These have been seen in normal cultures and also in those from dwarf spores. Work on these spores is now in progress, and up to the present there has been no indication that they possess the functions predicated for the "micro-conidia" by Ames in Podospora (Pleurage) anserina (1, 2, 3). The perithecium initials, whether normal or abortive, early send out branches to form secondary mycelia, but there is no evidence that any of these can be described as a trichogyne.

It is interesting to note that Winter in 1873 figured an ascus with

two spores in Sordaria (Podospora) anserina (7).

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A SIMPLE METHOD OF PRESERVING AND MOUNTING SPECIMENS OF FUNGAL LESIONS ETC. FOR DEMONSTRATION

By N. C. PRESTON

When engaged on infection experiments with Myrothecium roridum I had need of specimens of the lesions produced on stem and leaves which would remain fresh and demonstrable over a considerable period and which could be easily transported. The procedure here described was therefore adopted with considerable success and, since it is extremely simple and is very rapidly carried out, it is thought that this brief description will be of interest to other workers.

A 2 per cent. solution of agar containing about o 1 per cent. mercuric chloride is prepared and poured into Petri dishes of suitable size. When this has cooled to near solidifying point the leaves or other pieces of tissue are plunged, without previous preparation, directly into the agar and held beneath the surface with a warm needle until the mass has solidified sufficiently to retain them in position. The depth of agar in the dish will of course depend upon the thickness of the object to be embedded and should be just sufficient to cover it evenly. The agar must not be used too hot or the colour of the material will be lost.

Specimens prepared in this way will keep for many months, retaining their natural appearance even after the agar has become dry and hard. The method of mounting also has the advantage that both sides of the object can be closely examined since the lid of the dish may be safely removed without fear of mould contaminations. In specimens kept unsealed for six months hyphae and spores of the sporodochia present upon the leaves were still easily visible under

a $\frac{1}{6}$ in. objective.

NOTES ON TYPE SPECIMENS OF BRITISH INOPERCULATE DISCOMYCETES

(FIRST PART, NOTES 1-50)

By J. A. NANNFELDT (Botaniska Institutionen, Upsala)

A GRANT from the University of Upsala (C. F. Liljewalch's resestipendium) made it possible for me to visit London in the summer of 1932 in order to study Discomycetes in the herbaria of the British Museum (Natural History) and of the Royal Botanic Gardens, Kew. When comparing a British fungus flora with one from the Continent one is struck by the number of species described from Britain and not known outside it. After studying the types of those British species I was able to establish the fact that most of them have been known for a long time on the Continent also, though under different names.

My main interest was devoted to the inoperculates, and in these notes the results of my examination of fifty British species of that group are given. The preparation of the list has been much delayed for various reasons, but I hope to be able to continue it soon. My work in London was greatly facilitated by the courtesy shown to me by the officials, and I wish to expense my profound gratitude to them all.

In this paper the species are arranged alphabetically, according to the names in Ramsbottom's List of British Discomycetes (Trans. Brit. mycol. Soc. IV). For the convenience of the reader, the synonymy and citations to the British fungus floras as well as to Saccardo's Sylloge Fungorum, Rehm's Discomyceten, and Boudier's Discomycetes d'Europe are appended. For further information references are frequently given to v. Höhnel's numerous papers (his "Fragmente zur Mycologie" being cited as "Fr. z. M." and his "Mycologische Fragmente" as "M. Fr.") and to my own work: "Studien über die Morphologie und Systematik der nicht-lichenisierten inoperculaten Discomyceten" in Nova Acta Reg. Soc. Sci. upsal. Ser. IV, vol. VIII, No. 2 (cited as Nannfeldt, 1932).

1. Belonidium Jerdoni (Cke. & Phill.) Massee, Brit. F. Fl. IV, 229. Peziza Jerdoni Cke. & Phill. in sched.

This name, as well as *Hyalinia incarnata* (Cke.) Boud. (note 15), is a synonym for *Pseudohelotium pineti* (Batsch ex Fr.) Fuck. This gathering was listed by Phillips as *Mollisia lurida* (Pers.) Phill. and declared to

be quite distinct from *Peziza pineti* Batsch. Phillips based his identification on a Fries specimen at Kew, labelled *P. lurida*. Massee took up a herbarium name by Cooke & Phillips for the fungus in question, stating that it is different from *P. pineti* Batsch = *P. lurida* Pers. I have studied the type of *P. Jerdoni* as well as the Fries specimen. They are both typical *Pseudohelotium pineti* as understood by v. Höhnel (Fr. z. M. No. 1224) who points out the very large variability in the shape and size of the spores of this species.

No specimen of *Peziza lurida* is to be found in Herb. Persoon (at Leyden), but the description leaves no doubt that he had the same

fungus in mind.

2. Belonium Arctii (Phill.) Sacc. Syll. F. VIII, 495 (No. 2044); Boud. Discom. d'Eur. p. 118.

Peziza (Mollisia) Arctii Phill. ap. Buckn. in Proc. Bristol Nat. Soc.

N.S. IV, 58.

Mollisia Arctii Phill. Brit. Discom. p. 183.

Belonidium Arctii Massee, Brit. F. Fl. IV, 225.

Pyrenopeziza Arctii Nannf. in Nova Acta Reg. Soc. Sci. upsal. Ser. IV, vol. VIII, No. 2, p. 142.

The type specimen, as well as the specimens distributed in Vize, *Micro. f. brit.* No. 476, are identical with the fungus I recently published as *Pyrenopeziza Arctii* (Phill.) Nannf. Until then it was recorded only from the type locality, but I have found it to be fairly common in Sweden.

3. Belonium filisporum (Cke.) Sacc. Syll. F. VIII, 494 (No. 2039); Boud. Discom. d'Eur. p. 118.

Peziza (Mollisia) filispora Cke. in Grev. III, 66.

Belonidium filisporum Phill. Brit. Discom. p. 152; Massee, Brit. F. Fl. IV, 226.

Trichobelonium filisporum Rehm in Ber. bayer. bot. Ges. XIV, 107.

This species is said in the original description to grow "on sheaths of grass". The type specimen (in Herb. Kew.) shows that the grass is *Brachypodium sylvaticum* and the fungus is identical with *Belonopsis pallens* (Sacc.) Keissl. (For description and synonymy of this species see Nannfeldt, 1932, p. 104.) As Cooke's specific epithet antedates that of Saccardo, the fungus should be called *Belonopsis filispora* (Cke.) Nannf. n.comb.

The same species is preserved in Phillips's herbarium (Brit. Mus.) as "Belonidium albido-virella n.s. ined."

4. Calloria coniicola Cke. & Phill. ap. Phill. Brit. Discom. p. 333; Sacc. Syll. F. viii, 639 (No. 2637); Massee, Brit. F. Fl. IV, 152; Boud. Discom. d'Eur. p. 101. Judging from descriptions only, I have suggested that this species might belong to the genus *Laetinaevia* Nannf. (Nannfeldt, 1932, p. 191). The examination of the type specimen showed that the descriptions are very insufficient and that my suggestion was wrong. It is a very delicate *Lachnum*, which—so far as I understand—cannot be specifically separated from *L. brevipilum* v. Höhn.

5. Calloria cornea (B. & Br.) Phill. Brit. Discom. p. 332; Sacc. Syll. F. viii, 640 (No. 2639); Massee, Brit. F. Fl. iv, 152; Boud. Discom. d'Eur. p. 101.

Peziza cornea B. & Br. in Ann. Mag. nat. Hist. (2), VII, 183 (No. 578); Berk. Outl. p. 371; Cke. Handb. p. 704 (No. 2113).

This species has been studied by v. Höhnel (e.g. Fr. z. M. No. 1074), who considered it to be a *Mollisia*, *M. cornea* (B. & Br.) v. Höhn. I supported this view (Nannfeldt, 1932, p. 126), and gave drawings of the excipulum. v. Höhnel's opinion—as well as mine—was based solely on the study of specimens distributed in Rabenhorst's *Fungi europaei*. The type specimen is identical with these.

6. Calloria fusarioides (Berk.) Fr. S. Veg. Scand. p. 359; Phill. Brit. Discom. p. 331; Rehm, Discom. p. 463; Sacc. Syll. F. VIII, 639 (No. 2634); Massee, Brit. F. Fl. IV, 151; Boud. Discom. d'Eur. p. 101.

Peziza fusarioides Berk. in Mag. 700l. Bot. 1, 46 (No. 12); Berk. Outl. p. 371; Cke. Handb. p. 704 (No. 2114).

This species has always been interpreted correctly. A very full description is given by v. Höhnel (Fr. z. M. Nos. 1063-4).

7. Calycella claroflava (Grev.) Boud. Discom. d'Eur. p. 95.

Peziza claroflava Grev. Fl. Edin. p. 424; Berk. in Engl. Fl. v, pt. 2,
p. 203.

Helotium claroflavum Berk. Outl. p. 372; Cke. Handb. p. 713 (No. 2150); Phill. Brit. Discom. p. 165; Sacc. Syll. F. VIII, 225 (No. 914); Massee, Brit. F. Fl. IV, 233.

In Herb. Kew. I saw what may be the specimen referred to by Massee, viz. one labelled in Cooke's handwriting: "Helotium claro-flavum—ex Grev." This specimen, as well as numerous others both at Kew and in Brit. Mus., is typical young Calycella citrina (Hedw. ex Fr.) (= Helotium citrinum Fr.). In this connection the following note by Phillips is of a certain interest: "Most of my British specimens are immature" (in litt. ad Cooke, 18. v. 1876). The immaturity of the apothecia is evidently the cause of the small size of the spores.

8. Calycella flava (Klotzsch) Boud. Discom. d'Eur. p. 95.

Peziza flava Klotzsch in sched.

Helotium flavum Phill. Brit. Discom. p. 156; Sacc. Syll. F. VIII, 225

(No. 915); Massee, Brit. F. Fl. IV, 241.

The type specimen, which was very fully described by Massee, in my opinion represents typical *Calycella citrina* (Hedw. ex Fr.). It differs only externally by the somewhat horny, semitranslucent apothecia of a more uniform yellowish white. In winter, after frost, the apothecia of *C. citrina* undergo, however, just these changes. As to anatomical structure, asci and spores no differences could be detected.

9. Cudoniella Allenii A. L. Sm. in Trans. Brit. mycol. Soc. III, 40;

Sacc. Syll. F. xxII, 603 (No. 5331).

The type specimen in Herb. Mus. Brit. is typical Corynella atrovirens (Pers.) Boud. The apothecia are remarkably well developed with convex hymenium, and their stemlike bases are somewhat more conspicuous than usual.

It will be shown below that *Pachydisca agaricina* (Carm. ap. Berk.) Boud. (note 30) and *Pithyella hydnicola* (B. & Br.) Boud. (note 30)

are also Corynella atrovirens.

10. Dasyscypha campylotricha A. L. Sm. in Trans. Brit. mycol. Soc. III, 112; Sacc. Syll. F. XII, 684 (No. 5616) ("campylotrichia").

This species belongs to the genus *Unguiculella* v. Höhn. The specimens distributed by Karsten in F. Fenn. No. 652 (on *Artemisia vulgaris*) as *Peziza eurotioides* Karst. are the same fungus. It is evidently very rare, collected only in the type gatherings.* Its correct name is *Unguiculella eurotioides* (Karst.) Nannf. n.comb.

11. Dasyscypha crucifera (Phill.) Sacc. Syll. F. VIII, 440 (No. 1833); Massee, Brit. F. Fl. IV, 331; Boud, Discom. d'Eur. p. 119.

Peziza crucifera Phill. in Gard. Chron. (1878), p. 397. Lachnella crucifera Phill. Brit. Discom. p. 250.

This species belongs—as the descriptions also bear witness—to the genus *Lachnum Karst.*, and its correct name is *Lachnum cruciferum* (Phill.) Nannf. n.comb.

As far as I am aware, this fungus has never been reported from outside Britain, but I have seen it several times in Sweden and think

it is common everywhere on Myrica Gale.

Discinella exidiiformis (B. & Br.) Boud. Discom. d'Eur. p. 96.
 Peziza exidiiformis B. & Br. in Ann. Mag. nat. Hist. (4), xv, 37 (No. 1480); Cke. Mycogr. f. 60; Phill. Brit. Discom. p. 81; Massee in J. linn. Soc. (Bot.), xxxi, 501.

^{*} My report of *Peziza eurotioides* from Sweden (*Svensk bot. Tidskr.* xxII, 131 as *Pezizella eurotioides*) is based upon an erroneous determination.

Humaria exidiiformis Sacc. Syll. F. VIII, 122 (No. 468); Massee, Brit. F. Fl. IV, 418.

This species belongs to the operculate Discomycetes, probably to the genus *Humaria* (= *Humarina* Seav.), as Saccardo and Massee (*Brit. F. Fl.*) suggested.

13. Encoelia Bloxami (Berk.) Phill. Brit. Discom. p. 338; Boud. Discom. d'Eur. p. 161.

Patellaria Bloxami Berk. in sched.

Cenangium Bloxami Sacc. Syll. F. VIII, 568 (No. 2343); Massee, Brit. F. Fl. IV, 114.

As noted already by Massee on the type specimen (at Kew), this species is identical with Diplocarpa Curreyana Massee.

14. Habrostictis lasia (B. & Br.) Boud. Discom. d'Eur. p. 102.

Peziza lasia B. & Br. in Ann. Mag. nat. Hist. (4), II, 347 (No. 1391). Calloria lasia Phill. Brit. Discom. p. 327.

Orbilia lasia Sacc. Syll. F. VIII, 625 (No. 2574); Rehm, Discom. p. 456; Massee, Brit. F. Fl. IV, 146.

This species was treated earlier both by v. Höhnel (Fr. z. M. No. 1016) and myself (Nannfeldt, 1932, p. 97), but we did not study the actual type. The type specimen proves to be identical with the species treated by us and generally known as *Habrostictis lasia* (B. & Br.) Boud. or *H. rubra* Fuck. The latter is the valid name.

15. Hyalinia incarnata (Cke.) Boud. Discom. d'Eur. p. 103.

Peziza (Mollisia) incarnata Cke. in Grev. 1, 131.

Mollisià incarnata Phill. Brit. Discom. p. 191; Massee, Brit. F. Fl. 1V, 216.

Pezizella incarnata Sacc. Syll. F. VIII, 285 (No. 1186).

The type specimen (in Herb. Kew.) is in a rather bad condition and very poor, but it allows of the certain identification of this species as *Pseudohelotium pineti* (Batsch ex Fr.) Fuck. Further notes on this species are given above under *Belonidium Jerdoni* (Cke. & Phill.) Massee (note 1).

16. Lachnella canescens (Cke.) Phill. Brit. Discom. p. 259; Sacc. Syll. F. VIII, 394 (No. 1620); Boud. Discom. d'Eur. p. 123.

Peziza canescens Cke. in litt. ad Phill.

Dasyscypha canescens Massee, Brit. F. Fl. IV, 346.

Phillips placed this species next to Lachnella corticalis Fr. but considered it specifically distinct because of its somewhat different spores and "the more conspicuous septate hairs of the exterior". Examination of the type specimen (at Kew) proved that the alleged differences are very slight, if any, and fall within the variation shown by L. corticalis.

I have shown (Nannfeldt, 1932, p. 265) that the generic name Lachnella cannot be used in this sense, and that the species is a Lachnum, L. corticale (Pers. ex Fr.) Nannf.

17. Lachnella Crosslandi Boud. ap. Ramsb. in Trans. Brit. mycol. Soc. IV, 375.

Echinella Crosslandi Massee, Brit. F. Fl. IV, 306.

Pirottaea Crosslandi Sacc. in Hedw. xxxv, Beih. 7, p. xxxvi; Sacc. Syll. F. xxv, 776 (No. 2913).

The type specimen (at Kew) is Lachnum corticale (Pers. ex Fr.) Nannf. (see Nannfeldt, 1932, pp. 129 and 265) with unusually well-developed spores.

18. Lachnella (Helotiella) Laburni Phill. in Scot. Nat. (1891), p. 90; A. L. Smith in Trans. Brit. mycol. Soc. 1v, 76.

Helotiella Laburni "A. L. Sm." Sacc. Syll. F. XXIV, 1209 (No. 7267).

The type specimen (at Brit. Mus.) is very poor. The only Discomycete that I could find on it was *Unguicularia scrupulosa* (Karst.) v. Höhn. Phillips's description matches it tolerably, except for the asci and the spores. It is impossible to tell whether he made mistakes when measuring them or studied a mixture of two species.

19. Lachnella setulosa (Massee & Crossl.) Boud. ap. Ramsb. in Trans. Brit. mycol. Soc. IV, 375.

Echinella setulosa Massee & Crossl. ap. Massee, Brit. F. Fl. IV, 305. Pirottaea setulosa Sacc. in Hedw. xxxv, Beibl. 7, p. xxxvi; Sacc. Syll. F. xIV, 776 (No. 2912).

I have recently suggested that this species might be identical with Trichobelonium obscurum Rehm (Nannfeldt, 1932, p. 167). In Crossland's herbarium (now at Kew) there may be found not only the type collection but also seven additional gatherings. They are all identical with Rehm's species. T. obscurum Rehm is the valid name.

20. Lachnella siparia (B. & Br.) Phill. Brit. Discom. p. 276; Sacc. Syll. F. vIII, 396 (No. 1629); Boud. Discom. d'Eur. p. 123.

Peziza (Fibrina) siparia B. & Br. in Ann. Mag. nat. Hist. (2), XIII, 465 (No. 772); Berk. Outl. p. 370; Cke. Handb. p. 696 (No. 2079).

Dasyscypha siparia Massee, Brit. F. Fl. IV, 367.

The type (at Kew and Brit. Mus.) shows that *Peziza siparia* is a true *Encoelia* (Fr.) Karst. (see Nannfeldt, 1932, pp. 303-4) and identical with *Cenangium Ulmi* Tul. The British specimens are not—as Berkeley & Broome state—"on decorticated elm branches" but on

the inner bark of elm. As the specific epithet "siparia" antedates "Ulmi", the species should be known as Encoelia siparia (B. & Br.) Nannf. n.comb.

21. Lecanidion clavisporum (B. & Br.) Sacc. & D. Sacc. ap. Sacc. Syll. F. xvIII, 184 (No. 3808); Boud. Discom. d'Eur. p. 154.

Patellaria clavispora B. & Br. in Ann. Mag. nat. Hist. (2), XIII, 465 (No. 774); Berk. Outl. p. 373; Cke. Handb. p. 717 (No. 2166); Phill. Brit. Discom. p. 366; Massee, Brit. F. Fl. IV, 102; Massee in J. linn. Soc. (Bot.), XXXV, 107.

Durella clavispora Sacc. Syll. F. VIII, 794 (No. 3257).

Two authentic gatherings of this species were seen by me, viz. the type collection from Lucknam Grove (at Kew and Brit. Mus.) and a second gathering from St Catharine's (Herb. Broome in Brit. Mus.). They are both identical with *Lecanidion Crataegi* (Phill.) Sacc. (see next entry) and *Patellaria corticola* Starb.

I have published a photomicrograph of *P. corticola* and pointed out (Nannfeldt, 1932, p, 196) that it shows no relationship to the genus *Patellaria*. I know no other genus where it could be placed,

and its affinity is still obscure to me.

22. Lecanidion Crataegi (Phill.) Sacc. Syll. F. VIII, 799 (No. 3276); Boud. Discom. d'Eur. p. 155.

Patellaria Crataegi Phill. in Grev. xVII, 46; Massee, Brit. F. Fl. IV, 106.

In Herb. Phillips (at Brit. Mus.) are preserved two specimens under this name, both collected by Trail. The type specimen ("No. 3—on hawthorn twigs—Corbie Den—1. ii. 89") is identical with *Lecanidion clavisporum* (B. & Br.) Sacc. & D. Sacc. (see preceding entry). The second specimen ("No. 15—on bramble—nr Aberdeen—4. iii. 87") is a long-spored *Durella*.

The Romell specimen (Sweden: Gotland, Visby, 1. vii. 1887) in Herb. Kew. to which Massee (loc. cit.) alludes, is also Lecanidion

clavisporum.

23. Lecanidion Hyperici (Phill.) Sacc. Syll. F. VIII, 801 (No. 3288); Boud. Discom. d'Eur. p. 155.

Patellaria Hyperici Phill. in Grev. x, 69; Phill. Brit. Discom. p. 363; Massee, Brit. F. Fl. IV, 107.

This species, which is distributed in Phill. Elv. Brit. No. 191, is identical with Durella atrocyanea (Fr.) v. Höhn. (=Stictis atrocyanea Fr.). The apothecia are situated in greenish spots as they usually are in that species, though this feature was not mentioned in any description of Patellaria Hyperici. For further particulars about the very much misunderstood Durella atrocyanea see v. Höhnel, Ann. mycol., Berl., XVI, 210–12.

24. Lecanidion Lonicerae (Phill.) Sacc. Syll. F. VIII, 797 (No. 3267); Boud. Discom. d'Eur. p. 155.

Patellaria Lonicerae Phill. Brit. Discom. p. 364; Massee, Brit. F. Fl.

No specimen, but only a drawing was to be found in Phillips's herbarium (at Brit. Mus.). A specimen in Cooke's herbarium (at Kew), labelled by Phillips "Patellaria Lonicerae n.s.-On honeysuckle-Darnaway, N.B." is evidently part of the type material. The fungus is a true Durella, which I am unable to separate from D. vilis Starb. As Phillips's name is the older, the valid name of the species is D. Lonicerae (Phill.) Nannf. n.comb. (see Nannfeldt, 1932, p. 293).

25. Mollisia atrocinerea (Cke.) Phill. Brit. Discom. p. 176; Sacc. Syll. F. VIII, 322 (No. 1334); Massee, Brit. F. Fl. IV, 208; Boud. Discom. d'Eur. p. 136 (non Rehm, Discom. p. 530).

Peziza atrocinerea Cke. F. Brit. Exs. Ed. ii, no. 382.

As both Massee (loc. cit.) and Morgan (J. Mycol. vm, 182) have already suggested, this species, on Polygonum, is identical with the older Mollisia Polygoni (Lasch) Rehm. Rehm lists a M. atrocinerea (Cke.) Phill. which he regards as very closely allied to M. atrata (Pers. ex Fr.) Karst. and to M. revincta Karst. Rehm's species grows on stems of several different herbs but not on Polygonum. His herbarium shows that he placed numerous different fungi under this name as well as under Mollisia atrata.

M. Polygoni is a very characteristic species, differing in many respects from both Mollisia and Pyrenopeziza. Probably it should be placed in a separate genus of its own, but the time is not yet ripe

for a definite system of Mollisioideae.

26. Mollisia Browniana (Blox. ap. B. & Br.) Sacc. Syll. F. viii, 327 (No. 1355); Boud. Discom. d'Eur. p. 137.

Peziza (Mollisia) Browniana Blox. ap. B. & Br. in Ann. Mag. nat. Hist. (3), xv, 446 (No. 1072); Cke. Handb. p. 702 (No. 2102); Phill. Brit. Discom. p. 408.

Pseudopeziza Browniana Massee, Brit. F. Fl. IV, 199.

Examination of the type collection (at Kew and Brit. Mus.) proved that this species has been totally misinterpreted. The fungus is Heterosphaeria Patella Grev., and the substratum is not Epilobium hirsutum but some kind of Umbelliferous plant, almost certainly Angelica silvestris.

27. Niptera Stockii (Cke. & Phill.) Boud. Discom. d'Eur. p. 141. Peziza Stockii Cke. & Phill. in sched. Lachnella Stockii Phill. Brit. Discom. p. 261. Belonium Stockii Sacc. Syll. F. vIII, 496 (No. 2048). Echinella Stockii Massee, Brit. F. Fl. IV, 307.

The type specimen (at Kew) is in a very bad condition. It could be ascertained, however, that the fungus belongs to the genus *Pyreno-peziza* Fuck. emend. Nannf. It has septate spores and the margin of the excipulum passes into long, obtuse, hyaline hairs. The taxonomic position is next to *P. Arctii* (Phill.) Nannf. and *P. leucostoma* (Karst.) Nannf. Its identity must be left undecided until the host plant can be determined. It is probably a Composite.

28. Orbilia Boydii A. L. Sm. & Ramsb. in Trans. Brit. mycol. Soc. II, 168; Sacc. Syll. F. xxIV, 1239 (No. 7370).

The type specimen (in Herb. Brit. Mus.) as well as another authentic specimen ("on dead twigs of *Vaccinium Myrtillus*—Beith, Ayrshire—20th July 1912—D. A. Boyd") both show a very young, hardly ripe *Pezicula*, which can with certainty be identified with *P. myrtillina* Karst.

29. Orbilia scotica Massee, in Grev. XXII, 99; Massee, Brit. F. Fl. IV, 144; Sacc. Syll. F. XI, 426 (No. 2661); Boud. Discom. d'Eur. p. 103.

In the original description of this species Massee gives the information that it is based on a specimen from "Aboyne, N.B." lying in the Berkeley herbarium at Kew as *Peziza vinosa*. On a sheet of *Orbilia* vinosa there, I found a specimen marked in Berkeley's handwriting: "Pez. (Mollisia)—Aboyne, Sept. 1870." In my opinion this specimen must be the type, though it bears no note at all in Massee's hand. It matches the description of *Orbilia scotica* very well, except for the spores. These were extremely difficult to see, as they usually are in herbarium specimens of Orbilia, and the poorness of the material did not permit studying more than one apothecium. Nevertheless, I was able to find a few genuine free spores which were undoubtedly ascospores. They were needle-shaped and about $12 \times 1 - 1.5 \mu$. I have no hesitation in saying that Massee's description of the spores as "ellipticoblong, ends obtuse, $4 \times 1 \mu$ " was due to faulty observation. O. scotica is identical with the species that W. Nylander (Not. Sällsk. F. Fl. Fenn. Förh. x, 56) described as Peziza vinosa A. & S. and that has later been known as Orbilia vinosa (A. & S.) Karst. It is a typical Orbilia, but it is impossible to decide whether it is identical with the original Peziza vinosa of Albertini & Schweinitz.

30. Pachydisca agaricina (Carm. ap. Berk.) Boud. Discom. d'Eur. p. 93. Peziza agaricina Carm. ap. Berk. in Engl. Fl. v, pt. 2, p. 207. Helotium agaricinum Berk. Outl. p. 371; Cke. Handb. p. 708 (No. 2127); Sacc. Syll. F. vIII, 220 (No. 896). Belonidium agaricinum Massee, Brit. F. Fl. IV, 224.

The type specimen (in Herb. Kew.) proved to be Corynella atrovirens (Pers.) Boud.

31. Pachydisca brunnea (Phill.) Boud. Discom. d'Eur. p. 94.

Ombrophila brunnea Phill. in Grev. VIII, 103; Phill. Brit. Discom.
p. 323; Sacc. Syll. F. VIII, 619 (No. 2551); Massee, Brit. F. Fl.
IV, 143.

This species belongs to the operculates and probably to the genus *Humaria* (= *Humarina* Seav.), but nothing definite about its identity can be said for the present.

32. Pachydisca Laburni (B. & Br.) Boud. Discom. d'Eur. p. 94.

Helotium Laburni B. & Br. in Ann. Mag. nat. Hist. (4), XVII, 143

(No. 1624); Sacc. Syll. F. VIII, 249 (No. 1027); Massee, Brit.

F. Fl. IV, 235; Massee in J. linn. Soc. (Bot.), XXXI, 475.

Hymenoscypha Laburni Phill. Brit. Discom. p. 135.

Rehm (Discom. p. 787) identified this species with the older Helotium infarciens Ces. & deNot. This identification is correct.

33. Pachydisca ochracea (Grev.) Boud. Discom. d'Eur. p. 93.

Peziza ochracea Grev. Scott. Crypt. Fl. pl. 5; Berk. Engl. Fl. v, pt. 2, p. 204.

Helotium ochraceum Berk. Outl. p. 372; Cke. Handb. p. 713 (No. 2148); Phill. Brit. Discom. p. 169; Sacc. Syll. F. VIII, 229 (No. 937); Massee, Brit. F. Fl. IV, 237.

The type of this species is evidently lost, and Massee based his description on the original one, adding microscopical details from a Carmichael specimen (determined by Klotzsch) in Herb. Kew. This specimen is still there, but I could detect no Discomycete on it. If one may make a guess from Greville's drawing and description, I should suggest that they represent some species of *Pezicula*, tentatively *P. livida* (B. & Br.) Rehm. The description of the hymenium "as if sprinkled with minute shining particles not unlike small grains of brown sugar", strongly suggests that genus (see Nannfeldt, 1932, p. 90).

Massee's description of the microscopical features indicates that the Carmichael specimen also belonged to the genus *Pezicula*.

Specimens in Herb. Brit. Mus. from Broome's herbarium marked "Peziza ochracea—Hartham Park—21. iii. 43", are old Calycella citrina (Hedw. ex Fr.).

34. Pachydisca quisquilaris (Phill.) Boud. Discom. d'Eur. p. 94.

No such species was ever described, Boudier's reference "quisquilaris Phill., Grev. xvi, p. 94—Sacc. Syll. F. viii, p. 617" being a mixture of Helotium quisquiliaris Karst. and Ombrophila helotioides Phill.

35. Pachydisca scoparia (Cke.) Boud. Discom. d'Eur. p. 94.

Helotium scoparium Cke. in Grev. IV, III; Phill. Brit. Discom. p. 168; Sacc. Syll. F. vm, 239 (No. 974); Massee, Brit. F. Fl. IV, 234.

The type specimen at Kew represents a typical Pezicula Rubi (Lib.) Niessl, but the apothecia have (from age or bad preservation) lost most of their characteristic colour. It proved, on microscopical examination, to be unchanged internally.

36. Patinella Euphorbiae (B. & Br.) Sacc. Syll. F. vIII, 771 (No. 3171); Boud. Discom. d'Eur. p. 146.

Peziza (Patella) Euphorbiae B. & Br. in Ann. Mag. nat. Hist. (5), III, 212 (No. 1820).

Mollisia Euphorbiae Phill. Brit. Discom. p. 198.

Pseudopeziza Euphorbiae Massee, Brit. F. Fl. IV, 197.

I have studied the type specimen in Herb. Brit. Mus. The species proves to be identical with Naevia tithymalina (J. Kze.) Rehm (=Calloria tithymalina J. Kze.). The material is old and in bad condition; hence the dark colour.

Kunze's specific epithet antedates "Euphorbiae" by three years. The correct taxonomic position of the fungus is not clear, though I believe it to be closely related to my genus Laetinaevia (see Nannfeldt, 1932, pp. 190-2).

37. Peristomialis Berkeleyi Boud. Discom. d'Eur. p. 116.

Peziza (Mollisia) peristomialis B. & Br. in Ann. Mag. nat. Hist. (3), xvIII, 126 (No. 1169); Cke. Handb. p. 706 (No. 2119); Massee in J. linn. Soc. (Bot.), xxxv, 99.

Mollisia peristomialis Phill. Brit. Discom. p. 201. Cyathicula peristomialis Sacc. Syll. F. VIII, 308 (No. 1284); Massee,

Brit. F. Fl. IV, 273 ("peristomalis").

This very curious species was the only member of the subgenus Peristomialis of Mollisia in Phillips's Brit. Discom., and Boudier later established the genus Peristomialis, changing the specific name of the species into P. Berkeleyi. The taxonomic position of the genus was discussed by v. Höhnel (M. Fr. p. clviii), and he suggested from the description and the illustrations that it might belong to the "Nectriaceae?. When studying the type specimen I recognised at once that v. Höhnel had been right in excluding the genus from the Discomycetes and that, in reality, it is identical with Ijuhya Starb. V. Höhnel has given several good descriptions of this hitherto monotypic genus (Denkschr. Akad. Wiss. Wien, LXXXIII, 22; Fr. z. M. pp. 691 and 762). I cannot for the present decide whether Peziza peristomialis is identical

with Ijuhya vitrea Starb., known from Brazil and Java, or specifically distinct from it.

38. **Phaeangium phaeosporum** (Cke.) Sacc. & Syd. ap. Sacc. Syll. F. xvi, 765; Boud. Discom. d'Eur. p. 162.

Cenangium phaeosporum Cke. in Grev. XII, 44; Phill. Brit. Discom. p. 346; Sacc. Syll. F. VIII, 570 (No. 2354).

Schweinitzia phaeospora Massee, Brit. F. Fl. IV, 135.

The type specimen in Herb. Kew. is poor and in a very bad condition. No character could be detected that would distinguish it from *Velutaria rufo-olivacea* (A. & S. ex Fr.) Fuck. except the more uniform darker colours, which are certainly due to bad preservation. Thus both the original species of *Schweinitzia* Massee (non Elliot) are *Velutaria rufo-olivacea* and Massee's genus becomes synonymous with *Velutaria* Fuck. (see Nannfeldt, 1932, p. 302).

Saccardo & Sydow placed Cenangium phaeosporum in the genus Phaeangium (Sacc.) Sacc. & Syd. (non Pat.). The first species of that genus, P. Rubi (Bäuml.) Sacc. & Syd., is most probably a synonym for Pezicula Rubi (Lib.) Niessl (see Nannfeldt, 1932, p. 92), and the sixth and last of the original species, Phaeangium patellatum (Cke.) Sacc. & Syd. (=Cenangium patellatum Cke.), described from U.S.A. as growing on branches of Acer, is, according to the type specimen (at Kew), Dermatea Cerasi Fr. on Prunus. It seems most probable that the remaining species of Phaeangium Sacc. & Syd. are stages of well-known species with spores coloured because of age or bad preservation.

The genus *Phaeangella* (Sacc.) Massee seems to be in a similar position. Two species described by Hazslinsky as *Cenangium quercinum* (on oak) and *Tympanis Potentillae* (on *Potentilla fruticosa*) and later transferred to *Phaeangella* are both according to authentic material at Kew *Dermatea Cerasi* Fr. (on *Prunus*!). It is almost incredible how careless mycologists have been in describing "new" species and in naming their host plants. *Phaeangella Prunastri* (Fr.) Massee and *P. Morthieri* (Fuck.) Sacc. & Syd. are also species of *Dermatea*.

39. Pithyella hydnicola (B. & Br.) Boud. Discom. d'Eur. p. 125.

Peziza (Mollisia) hydnicola B. & Br. in Ann. Mag. nat. Hist. (4),

VII, 434 (No. 1327).

Mollisia (Mollisiella) hydnicola Phill. Brit. Discom. p. 194. Pseudohelotium (Mollisiella) hydnicola Sacc. Syll. F. VIII, 304 (No. 1269) ("hydnicolum").

Mollisiella hydnicola Massee, Brit. F. Fl. IV, 223.

This species, which has remained very doubtful and known only from the very short original description, is represented in Broome's

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herbarium (at Brit. Mus.) by part of the type gathering. Very superficial inspection showed that it did not grow on a Hydnum, but through a resupinate Hydnum. Its substratum is some kind of frondiferous wood, most probably oak. The fungus is typical Corynella atrovirens (Pers.) Boud. The spore description by Berkeley & Broome is totally false. They probably never saw the spores but mistook large drops of oil or protoplasm for them.

40. **Pseudopeziza foecunda** (Phill.) Massee, Brit. F. Fl. IV, 200; Boud. Discom. d'Eur. p. 180.

Peziza (Mollisia) foecunda Phill. ap. Stevenson, Mycol. Scot. p. 326. Mollisia foecunda Phill. Brit. Discom. p. 189.

Pyrenopeziza foecunda Sacc. Syll. F. viii, 369 (No. 1523).

This species, which was distributed in Phill. Elv. brit. No. 184, is identical with Hysteropezizella subsessilis (Rehm) Nannf. (see Nannfeldt, 1932, p. 121, where the full synonymy is to be found). The substratum given as "Eleocharis", is Scirpus caespitosus. As Phillips's name antedates the other names the fungus should be known as Hysteropezizella foecunda (Phill.) Nannf. n.comb.

Mollisia scirpina Peck (on Scirpus caespitosus) is certainly identical,

but I know it only from the description.

41. Pyrenopeziza cyanites (Cke. & Phill.) Boud. Discom. d'Eur. p. 133.

Mollisia cyanites Cke. & Phill. ap. Phill. Brit. Discom. p. 176; Sacc. Syll. F. VIII, 351 (No. 1452).

Belonidium cyanites Massee, Brit. F. Fl. IV, 225.

The type specimen of this species (at Kew) is very poor. The substratum was originally given as "some herbaceous stem". I was therefore very surprised to recognise it as *Phragmites communis*. The fungus was so sparse that I had to refrain from making a microscopical study. The macroscopical features and Massee's very full description indicate *Tapesia Kneifii* (Wallr.) v. Höhn. (= T. retincola (Rabenh.) Karst.) (see v. Höhnel, Fr. z. M. p. 1223). It is most unfortunate that mycologists so often describe "new" species on such insufficient material and that they take so little care in naming the host plants!

Mollisia mediella Karst. may be the same species. The superficial Mollisioid Discomycetes on *Phragmites* are greatly in need of a critical revision. Cultural experiments are needed to show to what extent the development of a dark subiculum depends upon external con-

ditions.

42. Pyrenopeziza grisella (Cke. & Phill.) Boud. Discom. d'Eur. p. 134.

Peziza grisea Carm. in sched.

Lachnella grisella Cke. & Phill. ap. Phill. Brit. Discom. p. 260. Trichopeziza grisella Sacc. Syll. F. VIII, 413 (No. 1702). Dasyscypha Carmichaeli Massee, Brit. F. Fl. IV, 363.

This species, according to the type specimen at Kew, belongs to *Unguicularia*. The hairs and the hymenium, etc., are exactly those of *U. scrupulosa* (Karst.) v. Höhn., but the apothecia are slightly larger than usual and the excipulum is remarkably dark, which may be due to age. We may safely regard the two species as synonymous. Karsten's specific epithet is the valid one.

43. Scleroderris majuscula Cke. & Massee, in Grev. xx1, 73; Massee, Brit. F. Fl. p. 125; Sacc. Syll. F. x1, 425 (No. 2652); Boud. Discom. d'Eur. p. 164.

The type specimen (at Kew) is very poor, consisting of a single, detached apothecium. It is *Coryne sarcoides* (Jacq. ex Fr.) Chev.

44. Tapesia Johnstoni (Berk.) Phill. Brit. Discom. p. 282; Sacc. Syll. F. viii, 381 (No. 1570); Boud. Discom. d'Eur. p. 140.

Peziza Johnstoni Berk. in Ann. Mag. nat. Hist. (1), XIII, 17 (No. 313); Berk. Outl. p. 369; Cke. Handb. p. 695 (No. 2075); Massee in J. linn. Soc. (Bot.), XXXI, 515.

After examining the type material of Peziza Johnstoni, Massee (loc. cit.) identified it with Tapesia fusca (Pers. ex Fr.) Fuck. I was very surprised when I saw the material at Kew to find that Massee had been totally wrong, for Peziza Johnstoni is identical with the fungus known as Cenangella radulicola (Fuck.) Rehm (= Cenangium radulicola Fuck. = Dermatea radulicola Fuck.) growing, as always, on branches of birch infested with Eutypa aterrima (Fr.) v. Höhn. (= Radulum aterrimum Fr.). Berkeley's name antedates Fuckel's.

As I have not yet found appropriate material in quantity I have not been able to study the species in detail and cannot decide its

taxonomic position.

45. Trichoscypha calycina var. Trevelyani (Cke.) Boud. Discom. d'Eur. p. 125 ("Trevyliani").

Peziza calycina var. Trevelyani Cke. in Grev. III, 121.

Lachnella calycina var. Trevelyani Phill. Brit. Discom. p. 242.

Dasyscypha calycina var. Trevelyani Sacc. Syll. F. vIII, 438 (sub No. 1822); Massee, Brit. F. Fl. IV, 342.

The type specimen of this variety is *Trichoscyphella Willkommii* (Hart.) Nannf. (= Dasyscypha Willkommii (Hart.)). I could not find any spores larger than $24 \times 8\mu$, and the average size was $20 \times 7\mu$,

just as in typical *Trichoscyphella Willkommii*. The larger measurements given by Cooke must be due to some mistake.

46. Trichopeziza dematiicola (B. & Br.) Sacc. Syll. F. VIII, 414 (No. 1707); Boud. Discom. d'Eur. p. 131.

Peziza (Mollisia) dematiicola B. & Br. in Ann. Mag. nat. Hist. (3), xv, p. 446 (No. 1070); Cke. Handb. p. 705 (No. 2117).

Lachnella dematiicola Phill. Brit. Discom. p. 265.

Dasyscypha dematiicola Massee, Brit. F. Fl. IV, p. 364 (saltem p.p.). Dasyscypha dematiicola V. Höhn. in S.B. Akad. Wiss. Wien, CXVIII, Abt. 1, p. 884.

The type specimen (at Kew) was studied by v. Höhnel (Fr. z. M. p. 339), who identified it with his *Dasyscypha Heimerlii* v. Höhn. I have studied the type at Brit. Mus. and can only assert that it matches v. Höhnel's description of *D. Heimerlii* in all respects.

The systematic position of this species is somewhat obscure. The texture of the excipulum and the characteristic pointed hairs would place it in *Hyaloscypha* Boud. emend. Nannf. but the brown colour of the excipulum (the hairs and the margin excepted) distinguish it from the known species of that genus. Nevertheless, I think it best for the present to call it *Hyaloscypha dematiicola* (B. & Br.) Nannf. n.comb. (see Nannfeldt, 1932, p. 272).

47. **Trochila Buxi** Capron ap. Cke. *Handb*. p. 768 (No. 2315); Phill. *Brit. Discom*. p. 397; Sacc. *Syll. F.* vIII, 729 (No. 2991); Massee, *Brit. F. Fl.* IV, 61; Boud. *Discom. d'Eur.* p. 166.

The type specimen (Herb. Kew.) as well as two additional British specimens (Forden, leg. Vize, and Bungay, leg. Stock) are all Hyponectria Buxi (DC.) Sacc. A very full description of this species was given by Petrak (Ann. mycol. XXI, 303-6), who also discusses its taxonomic position. It may be noted that Laestadia Buxi (Fuck.) Sacc. is the same species (v. Höhnel, M. Fr. p. cci), as is also the fungus that Feltgen reported as Trochila Buxi Capron (v. Höhnel, S.B. Akad. Wiss. Wien, cxv, Abt. I, p. 1263).

48. Urceolella elaphines (B. & Br.) Boud. Discom. d'Eur. p. 129.

Peziza elaphines B. & Br. in Ann. Mag. nat. Hist. (4), VII, 434 (No. 1325); Massee in J. linn. Soc. (Bot.), xxxv, 90.

Mollisia elaphines Gill. Champ. Franc. Discom. p. 131; Phill. Brit. Discom. p. 179.

Pseudohelotium elaphines Sacc. Syll. F. vIII, 301 (No. 1257).

Dasyscypha elaphines Massee, Brit. F. Fl. IV, 366.

The identity of Peziza elaphines B. & Br. and Peziza scrupulosa Karst. (= Unguicularia scrupulosa (Karst.) v. Höhn.) was demonstrated by

v. Höhnel (Mitt. Bot. Inst. Techn. Hochsch. Wien, v). I can only confirm v. Höhnel's statement.

49. Urceolella leuconica (Cke.) Boud. Discom. d'Eur. p. 130.

Peziza leuconica Cke. in sched.

Lachnella leuconica Phill. Brit. Discom. p. 267.

Trichopeziza leuconica Sacc. Syll. F. VIII, 414 (No. 1709).

Dasyscypha leuconica Massee, Brit. F. Fl. IV, 334.

The type specimen (at Kew) shows that it belongs to the genus Hyaloscypha Boud. emend. Nannf. (Nannfeldt, 1932, pp. 272-3.). The substrate is coniferous wood, probably pine. It is distinct from all species known to me by the much longer hairs. As closely allied forms are described in Trichopeziza, Lachnella, Dasyscypha, Pezizella, and many other genera, it is impossible for the present to form a definite opinion as to the validity of the specific epithet, but the species may ad interim be designated as Hyaloscypha leuconica (Cke.) Nannf. n.comb.

50. Urceolella Stevensonii (B. & Br.) Boud. Discom. d'Eur. p. 130 ("Stephensoni").

Peziza (Mollisia) Stevensoni B. & Br. in Ann. Mag. nat. Hist. (4), xv, 38 (No. 1485).

Lachnella Stevensoni Phill. Brit. Discom. p. 235.

Dasyscypha Stevensoni Sacc. Syll. F. VIII, 454 (No. 1889); Massee, Brit. F. Fl. IV, 364; Boud. Discom. d'Eur. p. 122.

Quite twenty years ago v. Höhnel (M. Fr. p. x) described Dasyscypha resinifera v. Höhn., a species said to grow on old, prostrate, still hard coniferous logs, and to be very common in Lower Austria, and also to occur in Germany and Sweden. I was later able to show (Nannfeldt, 1932, p. 273) that this most characteristic species was identical with Pezizella atomaria Starb. and belonged to Hyaloscypha Boud. emend. Nannf. During my stay in England I found that the fungus was described still earlier by Berkeley & Broome as Peziza Stevensoni, and that the substratum of the type material was pinewood. The valid name of this species therefore becomes Hyaloscypha Stevensoni (B. & Br.) Nannf. n.comb.

A specimen in Broome's herbarium (Brit. Mus.) is by a slip of the pen labelled *Peziza Andersoni*, and the collector's name given as

J. Anderson.

A LEAF-SPOT DISEASE OF SWEET WILLIAM CAUSED BY HETEROSPORIUM ECHINULATUM

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(With 3 Text-figures)

Introduction

The diseased Sweet William plants (Dianthus barbatus L.) which were utilised in the present investigation were sent to the Plant Pathological Laboratory of the Department of Agriculture for

Scotland in May 1932.

The leaves of the plants were heavily infected, bearing numerous brownish patches covered with olive-green conidia. The fungus was identified as *Heterosporium echinulatum* (Berk.) Cooke, and it was associated with some small black bodies on several of the over-wintered leaves. On examination a few of these bodies were found to contain mature asci, so, in view of the fact that no perfect stage of *Heterosporium echinulatum* has yet been recorded, the present investigation was undertaken primarily in order to ascertain whether these perithecia represented the perfect stage in the life history of this fungus. In the course of the work, however, certain other observations were made which have also been included in the present paper.

HISTORICAL

The fungus was first described and named in 1870 by Berkeley in an illustrated account of a new disease on carnation leaves. He placed the causal fungus in the genus *Helminthosporium* as a new species *H. echinulatum*.

In 1873 Berkeley & Broome published a description of a disease on Sweet William caused by another new species of Helminthosporium, H. exasperatum. Four years later Cooke transferred H. echinulatum to the genus Heterosporium, and cited Helminthosporium exasperatum as a

synonym.

In 1881 Saccardo and Roumeguère described a fungus on *Dianthus barbatus* and named it *Heterosporium Dianthi*, but this name, together with *H. exasperatum*, was later cited by Saccardo (10) as a synonym of *H. echinulatum*.

In most references to the disease the parasite appears to have been confined to the leaves and stems of the hosts, but in 1890 Lindemuth

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referred to a case where the mycelium spread to the sepals and pre-

vented the opening of the flowers.

No mention is made of sclerotia in any of the earlier references to the disease, but in 1906 Cooke stated that numerous minute sclerotia are said to be formed in the dying leaves. Massee (8) also mentioned sclerotia, and recorded that they remain in a passive condition until the following season when they produce minute conidia. From this paucity of references to sclerotia in early records of the fungus it would seem that they are not of frequent occurrence.

CULTURAL STUDIES

Ascospores were removed from crushed perithecia and placed in hanging drops of sterile water in a moist chamber. They germinated in less than twenty-four hours, each cell giving rise to a germ tube which grew rapidly and produced a branching mycelium in two to three days. In five to six days conidiophores developed which cut off typical Heterosporium echinulatum conidia, thus establishing the relation between the perfect and imperfect stage of this fungus. Single ascospores planted on malt extract agar gave rise to a somewhat dome-shaped mass of mycelium, white above and olive green underneath, which produced typical H. echinulatum conidia in four to five

weeks when the colony was about 1-14 in. long.

Single conidia were also isolated from the original material and were found to give rise to two distinct types of culture on malt agar. One (type A) was exactly similar to that derived from single ascospores, the other (type B) differed in the time that elapsed before the formation of conidia, which here was only four to ten days. When conidia were produced the surface colour of the culture changed, becoming greyish green in type A and olive green in type B. This difference in colour appeared to be due to the development of large numbers of conidia on a small culture in type B whereas they were more scattered in type A. Conidia from these two types were repeatedly reisolated and transplanted but the distinction persisted on malt agar. If, however, sterilised carnation and Sweet William leaves were utilised instead of an agar medium, conidia were produced freely by both A and B types after ten to twelve days.

There appeared to be a great deal of variation in size between conidia from different cultures on the same medium and also between those on different media. Conidia from sterilised carnation leaf cultures, which were subcultures of the original malt agar cultures, were markedly larger than those of the parent cultures, and conidia from these leaf cultures put back on malt agar produced a mycelium which

also gave rise to large conidia (Table I).

In all the malt agar cultures, whether derived from conidia or

Table I. Measurements of conidia (in μ)

Source of conidia	Range	Mean
Original material	16-52 × 8-14	31.71×10.81
Malt agar culture (A type) old	$18-55 \times 8-13$	30·3 × 10·47
,, (B type) ,,	18-51 × 8-14	30.86 × 10.86
,, (A type) young	$16-45 \times 7-13$	27.53 × 10.09
,, (B type) ,,	20-50×6-13	31.66× 9.5
Sterilised carnation leaf culture	26-54×9-14	38·42×11·01
Subculture from carnation leaf on malt agar, 6 days old	29-53×9-14	39·93×12·37

ascospores, dark knotted masses of hyphae appeared after about five weeks. These were thought to be young perithecia, but they did not develop further although the cultures were kept for six months. On sterilised carnation and Sweet William leaves immature perithecia developed after twelve to fifteen days, the majority of them being formed on the surface furthest from the point of inoculation. They were most abundant towards the base of the tube where there was a greater percentage of water, and they developed particularly at points where the leaf was in contact with the glass. Perithecia in fifteen days old cultures measured from 60 to 116 μ in diameter, were black, roughly spherical, but differed from those of the original material in having a large number of black hyphae projecting from the walls. A short typical beak was present and the apical cells of the beak were colourless. On crushing them no asci were seen, and the contents were found to consist of a ball of hyaline cells and numerous oil globules.

In an attempt to obtain mature perithecia some conidia were planted on sterilised Sweet William leaves on December 6, 1932, and kept in the laboratory until January 10, 1933, when two of them were placed in an unheated greenhouse. Perithecia were taken from all the cultures and examined every fortnight, and asci were found in the greenhouse cultures in May 1933. The percentage of perithecia containing asci was small, as in the original material, and many of the asci were degenerate, but a few contained fully developed ascospores which agreed in every respect with those found on the original material. None of the perithecia in cultures kept in the laboratory developed asci, but the perithecia in seven to eight months old cultures which were kept damp grew very irregular in shape and the beak often became elongated (Fig. 1 b). In cultures which were allowed to dry the perithecia remained spherical and exactly resembled those of the original material.

DESCRIPTION OF FUNGUS

The conidial stage has been frequently described (1, 2, 3, 8), and it is therefore proposed to proceed at once to the description of the perfect stage.

The perithecia were black and irregularly spherical, $100-270\,\mu$ in diameter, and possessed short stout beaks $10-30\,\mu$ high (Fig. 1 a). The outer part of the walls consisted of two to three layers of dark brown cells, and the inner part of a few layers of colourless cells.

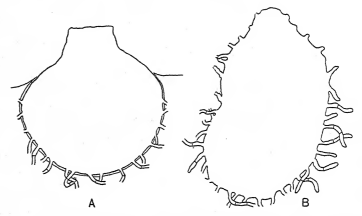


Fig. 1. Didymellina Dianthi. A, typical perithecium on original diseased material; B, abnormal perithecium from culture on Sweet William leaf. Perithecia × 400.

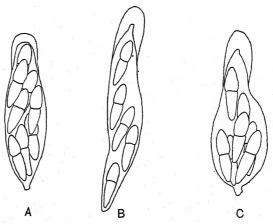


Fig. 2. Didymellina Dianthi. Asci from Sweet William leaf culture. A, typical ascus from centre of perithecium; B, abnormally elongated ascus; C, ascus from side of perithecium. (×500.)

The percentage of perithecia containing asci was small, about 10 per cent., and the remainder appeared to be sterile. When kept under moist conditions, however, many of the sterile perithecia developed tufts of conidiophores from the beak, the conidia exactly

resembling those of *Heterosporium echinulatum*. None of the fertile perithecia examined were producing conidia, so it would appear that sterile perithecia may function as sclerotia. This resembles the behaviour reported by Tisdale in the related species *H. gracile* (11).

The number of asci in a fertile perithecium varied from eight to eighteen according to the size of the perithecium. They were fascicled and attached by short pedicels to a mass of pseudoparenchyma in the base of the perithecium. Paraphyses were absent. The asci were thin-walled, hyaline and very irregular in shape. As the bundle of asci fitted exactly into the cavity of the perithecium, those towards the outside were shorter, stouter and club-shaped (Fig. 2 c), whereas those in the centre tended to be elongated and spindle-shaped (Fig. 2 a, b). The wall was thickened at the apex up to 10μ , and occasionally the cavity of the ascus was extended upwards into the centre of this thickening (Fig. 2 b, c).

Most of the asci contained eight ascospores lying parallel or slightly oblique to the long axis of the ascus. The ascospores measured $22-31 \times 7-9\mu$, and were torpedo-shaped, two-celled, thin-walled and colourless, the upper cell being $1-6\mu$ (average $3\cdot3\mu$) shorter, and slightly wider than the basal cell. The spores were somewhat con-

stricted at the septum.

INFECTION EXPERIMENTS

Infection experiments were carried out with pot plants of Sweet William, carnations and pinks. A suspension of conidia was injected into the leaves with a hypodermic syringe; the control plants were similarly treated, sterile water being used instead of the suspension. The conidia for some inoculations were obtained from monoascospore cultures, for others from type A or B cultures derived from single conidia off the original material. Ascospores were not utilised for these experiments owing to the scarcity of mature asci. The plants were kept under bell jars in the laboratory for forty-eight hours after inoculation and then placed in an unheated greenhouse and watered from below.

All the Sweet William plants developed lesions around the point of inoculation. These consisted of yellowish brown withered patches with a purplish margin (Fig. 3); they appeared in six to ten days and attained a diameter of about half an inch after four weeks. Very few conidia were found on these patches, so, since extremely dry conditions were prevalent in the greenhouse, an infected leaf was removed and placed on moist filter paper in a sterile Petri dish. Numerous hyphae developed within twenty-four hours on both sides of the infected area, and it was green with conidia forty-eight hours after removal to moist conditions. One of the infected plants was then placed under a bell jar, and it developed numerous conidia on the

infected areas after forty-eight hours. Conidia from these leaves were isolated on malt agar slopes and gave rise to typical cultures. The uninfected leaves of the plants remained healthy and the plants flowered.

Two months after inoculation the diseased areas had extended over the greater part of the infected leaves, but the plants were otherwise healthy. The control plants remained healthy and the injured points healed leaving only a small yellow scar.

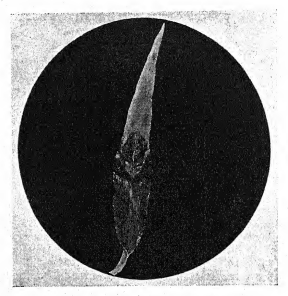


Fig. 3. Sweet William leaf artificially injected with *Heterosporium echinulatum* photographed two months after inoculation. The upper half of the leaf has been killed and the diseased area is delimited by a dark purple margin. About two-thirds natural size.

The injected carnations and pinks, on the other hand, showed only a very slight reaction, consisting of a yellowish area with a definite purple margin which, after four weeks, extended about $\frac{1}{8}$ in. around the point of inoculation. No Heterosporium conidia were found on the infected leaves, but in one or two sections hyphae were found on the injured part of the epidermis apparently growing saprophytically. Two leaves of pink and two of carnation were therefore cut off and kept moist in sterile Petri dishes. In two days hyphae developed on both of the pink leaves and on one of the carnation leaves, and conidia appeared in three days. These were Cladosporium on both pink leaves and on one of the carnations, but on the other carnation a few Heterosporium echinulatum conidia were also found. The mycelium,

however, did not spread to the uninjured part of the leaf which

remained free from hyphae.

The wound made by the needle on the controls healed, leaving a small yellow withered piece of epidermis, but no extension of the necrotic area was observed and no purple discoloration developed.

Table II. Infection experiments

		Total no. of						
		leaves	Re	sult	No. of	No. of	Res	sult
	No. of	inocu-			control	leaves		
Host	plants	lated	Diseased	Healthy	plants	injected	Diseased	Healthy
Sweet William	3	22	20	2	I	6	О	6
Carnation	ĕ	36	32*	4	ı	6	0	6
Pink	3	24	20*	$\dot{\tilde{4}}$	1	6	0	6
			* Slightl	y discolou	red.			

The results of the infection experiments are summarised in Table II. These inoculations were all carried out in early summer, but a subsequent series made on Sweet William in the autumn showed that infection did not take place so readily at that season,

neither did the lesions attain so great a size.

It is evident that the strain of *Heterosporium echinulatum* utilised in the present investigation is actively parasitic on Sweet William but has little effect on carnations and pinks. The two last-named plants, however, are well known as hosts of this fungus, a fact which leads to the conclusion that there must be two or more specialised races of the fungus upon the different host plants.

DISCUSSION

There appears to be no previous record of the perfect stage of Heterosporium echinulatum, nor could I find a description of any perithecial form on Sweet William, carnation or pink which corresponds to the perithecial stage described in the present paper. Tisdale (11) and Klebahn (6) have both published accounts of the perfect stage of H. gracile, the former identifying it as Didymellina Iridis (Desm.) v. H., while the latter regarded it as a new species which he named D. macrospora. The genus Didymellina was established by von Höhnel (5), but no adequate generic description was published. This fact somewhat complicates the identification of the perfect stage of Heterosporium echinulatum, but as the fungus agrees in essential points with the species described by Tisdale, it is thought advisable to use the name Didymellina for the perithecial form described in the foregoing pages. The specific epithet echinulatum refers to a character of the conidium wall which is by no means always well marked and which, in any case, is not applicable to the ascospores. The name Didymellina

Dianthi is therefore proposed for the new perfect stage of the fungus, since it occurs only, so far as we know, upon certain species of Dianthus.

Didymellina Dianthi n.sp.

Perithecia suberumpentia, atrobrunnea vel nigra, laxe gregaria, irregulariter sphaerica, $100-270\,\mu$ diametro, rostro brevi $10-30\,\mu$ longo. Asci in perithecio 8–18-nati, fasciculati, breviter pedicellati, quoad formam atque magnitudinem valde variabiles, muro hyalino infra tenui apicem versus ad $10\,\mu$ crasso, sporis octonis; paraphyses desunt. Ascosporae hyalinae, ovali-ellipticae, $22-31\times7-9\,\mu$, uniseptatae, ad septum paulo constrictae, cellula superiore paululo breviore atque latiore quam inferiore.

Hab. in foliis Dianthi barbati tempore hiberno emarcidis. Apud

hortos nonnullos in Scotia.

Forma conidialis—Heterosporium echinulatum (Berk.) Cooke.

SUMMARY

- 1. The perfect stage of *Heterosporium echinulatum* was found on some over-wintered leaves of diseased Sweet William plants, and was subsequently obtained in culture on sterilised Sweet William leaves. The relation between the two stages was proved by monospore cultures.
- 2. Only a small percentage of the perithecia matured; the remainder developed tufts of conidiophores from the beak if kept under moist conditions.
- 3. Infection experiments indicated the probability of specialised races of the fungus on different hosts, for Sweet William plants were very susceptible while carnations and pinks proved highly resistant.
- 4. The name Didymellina Dianthi is proposed for the new perfect stage, a diagnosis of which is given.

ACKNOWLEDGMENTS

My thanks are due to Prof. Sir William Wright Smith for granting facilities for this work at the Royal Botanic Garden, Edinburgh. I also desire to thank Mrs N. L. Alcock and Dr M. J. F. Gregor for obtaining material and for their advice and assistance throughout the investigation. I am indebted to Dr C. E. Foister for the photograph reproduced as Fig. 3, and to Mr Beaumont of Seale-Hayne Agricultural College for specimens of *Heterosporium gracile*. Miss E. M. Wakefield has kindly read and criticised the manuscript of this paper.

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CORDYCEPS MILITARIS AND ISARIA FARINOSA

By T. PETCH

In Ann. Sci. nat. Ser. 3, xx (1853), 43, Tulasne stated that he had reason to believe that the conidial stages of Cordyceps militaris, C. entomorrhiza, etc., were included in the genus Isaria as autonomous fungi. Subsequently, in the same journal, Ser. 4, VIII (1857), 35–43, he published the results of experiments which, he considered, proved that Isaria farinosa was the conidial stage of Cordyceps militaris. Figures in support of that conclusion were given in Selecta Fungorum Carpologia,

m (1865), Pl. I.

Tulasne obtained a number of dead or dying larvae of $Bombyx \ rubi$, the victims of a breeding experiment, and placed them on damp sand. They developed a white covering of mycelium, from which arose erect conidiophores, with whorls of branches, each terminated by a chain of conidia. He germinated these conidia in a hanging drop, and found that they reproduced the same conidiophores and conidia. Evidently that was the simple Spicaria form of $Isaria \ farinosa$, though Tulasne described the conidia as spherical, $I \cdot 5 \mu$ diameter, and rather confused matters by a reference to $Botrytis \ Bassiana$. His figures, however, show that he was not dealing with the latter species, and his description of the spores as spherical was perhaps due to the inadequacy of his lenses.

The byssoid covering soon became orange-yellow here and there, and developed *Isaria farinosa*. There is little doubt that Tulasne's identification of that was correct. Subsequently, many of the *Isaria*

clavae were attacked by Melanospora parasitica.

The experiment was begun in the middle of March. At the beginning of June, some of the larvae, which had produced only a few Isaria clavae, or none at all, their segments having remained only more or less whitened by the conidiferous mycelium, began to develop Cordyceps militaris, which matured at the beginning of July. Some of these Cordyceps clavae bore conidiophores towards the base.

In Selecta Fungorum Carpologia, III (1865), 6, Tulasne gave further details. He said that the conidiophores and conidia were the same, whether they occurred on the mycelium or on the Isaria clavae, and that they were very often to be found also on the "roots" or lowest parts of the Cordyceps clava. On germination of the ascospores (partspores) of the Cordyceps, hyphae were produced, from which arose branches bearing chains of conidia or even whorls of chains of conidia.

The conidia produced from the ascospores were exactly similar to

those of Isaria farinosa.

Tulasne's figures, Pl. I, figs. 19 and 20, show larvae bearing Isaria farinosa only. Figs. 21, 22, 23 show germinating conidia of I. farinosa, producing whorls of phialides bearing apical chains of conidia, evidently a regular Spicaria. Figs. 24, 25, 26 show larvae bearing Cordyceps militaris only, not accompanied by Isaria farinosa. Fig. 27 is that of the mycelium and conidiophores found at the base of a Cordyceps clava, and it shows one, two, or three phialides on the apex of a hypha and others below, but the latter are scattered, not in whorls; only one conidium is present at the apex of a phialide, but the absence of chains might be attributable to loss in manipulation. Fig. 28 is a view of the perithecial clava enlarged. Figs. 29 and 30 show germinating part-spores, and Fig. 31 shows a resultant hypha with conidiophores; the phialides are scattered along the hypha, or three occur on the apex of a short lateral branch; they bear chains of conidia, except in one case where three conidia are shown side by side at the apex of a phialide.

Judging these figures in the light of recent cultures of *Isaria farinosa* and of the ascospores of *Cordyceps militaris*, which will be described later in this paper, it is clear that while Figs. 27 and 31 represent, somewhat imperfectly, the conidial stage of *C. militaris*, they are not figures of *Isaria farinosa*. The fungus shown is not the regular *Spicaria* of *Isaria farinosa*. Other differences will be enumerated in the fol-

lowing pages.

Evidently, as might be expected, Tulasne's material contained a mixture of fungi. Some of the larvae were infected by *Isaria farinosa*, and these produced, first a white mycelium which bore simple *Spicaria* conidiophores, and later the clavae, of *Isaria* (*Spicaria*) farinosa. Others were infected with *Cordyceps militaris*, and, as the conidial stage of that fungus is not luxuriant in nature, they bore only a slight covering of conidiiferous mycelium, and ultimately produced *Cordyceps* clavae.

It may be mentioned that, from a collection of pupae of the Cinnabar moth, which had died during a breeding experiment at Farnham Royal, I obtained *Isaria farinosa*, *Beauveria Bassiana*, *Monilia penicillioides*, *Gymnoascus Reesii*, and undetermined species of *Fusarium*

and Stilbella.

Tulasne's theory met with considerable opposition. De Bary, after carrying out experiments on the subject, at first rejected it, but subsequently accepted it. The following details, taken from de Bary's Vergleichende Morphologie und Biologie der Pilze, etc. (1884), give the reason for his change of opinion.

"In [Bot. Zeitung, 1869] and 1867, I had expressed doubts concerning the view maintained by Tulasne that Isaria farinosa belonged to the cycle of development of C. militaris, basing those doubts partly

on the failure to obtain perithecial clavae and Isaria forms reciprocally from one another in culture, and partly on differences, certainly only quantitative, in the branching of the conidiophore. The latter objection might easily be dismissed, and, as already remarked in the foregoing text, I now believe that the former cannot be maintained. A caterpillar of Sphinx euphorbiae, which had been infected with ascospores and had become, as usual, a sclerotium, when laid on moist sand produced first two small perithecial clavae with normal perithecia. These died before the asci were fully developed, and then Isaria was produced in abundance. Portions of the mycelium cultivated on microscope slides had previously afforded Isaria. In this case, therefore, either Isaria was ultimately produced from the ascospores, or the insect had been infected with Isaria unintentionally at the same time as with the ascospores, and the Isaria had, in its later development, suppressed and supplanted the perithecial form. I have no reason for assuming such an accidental infection, and have accordingly formed the foregoing conclusion. But the possibility of such an admixture is not excluded, and therefore I could not omit to mention it."

De Bary stated that if the ascospores were sown in water or in a nutrient solution, germination occurred, with branching of the resultant hyphae dependent upon the amount of nutrient present. In water, only short hyphae, with few or no branches, were produced. Some of the branches spread through the nutrient solution as a mycelium; others emerged from the liquid into the air, where they produced whorls of phialides which bore conidia in chains. The first conidium on a phialide was cylindric, like those in the body of the insect, but usually shorter. All the succeeding conidia were globose. The latter could therefore be called globose or aerial conidia. The mycelium which developed in the dead caterpillar very often produced aerial conidia, but no cylindric conidia. The conidiophores which were found on most of the caterpillars which bore perithecial clavae were small, like those described above, and formed a delicate down on the surface. [On the other hand, on other insects they grew into a dense mould-like covering some millimetres in height, or, like the Coremium form of Penicillium, they formed clavate structures, 1-2 cm. high, with an orange-yellow stalk, covered above with phialides bearing conidia. These last-named bodies were known as a form species under the name of Isaria farinosa. Both the Isaria form and the mould-covering were usually found alone on the sclerotioid insect, without any perithecial clava.] Only once had he been able to obtain, on a caterpillar which had been infected with ascospores and which pupated after infection, two poorly developed perithecial clavae together with large Isariae.

Later in the same account (p. 401) de Bary stated that on insects killed by infection with aerial conidia, perithecial stromata were

never observed, but only a fresh crop of aerial conidia, especially the Isaria form.

In the foregoing summary, I have bracketed a section which is obviously a general account of *I. farinosa*, as it occurs in nature, not a series of observations by de Bary from his experiments, and which, as it happens, has no relation to the statements which precede it. Attention may be specially directed to de Bary's statement that the conidiophores which accompanied the *Cordyceps* clavae formed only a delicate down on the surface of the larvae.

De Bary identified as *Isaria farinosa* the conidiophores which he obtained on the germination of the ascospores of *Cordyceps militaris* and those found on caterpillars which bore perithecial clavae of that species. Probably for that reason it is not always clear from his account whether in his infection experiments with "aerial conidia" he made use of conidia obtained by germination of the ascospores or conidia taken from naturally grown specimens of *Isaria farinosa*. For example, in the statement quoted above from p. 401, aerial conidia would appear to mean conidia taken from *I. farinosa*. Nor is it always certain whether his references to "*Isaria*" mean the compound Isarioid form or the individual conidiophores, *e.g.* in his statement that "*Isaria*" developed from mycelium on microscope slides.

It will be noted that de Bary observed differences between the branching of the conidiophores obtained by germination of the ascospores of Cordyceps militaris and that of the conidiophores of Isaria farinosa, but did not, finally, consider them of importance. Both Tulasne and de Bary appear to have restricted their comparisons to conidiophores obtained by germination of the ascospores of the Cordyceps and of the conidia of the Isaria, respectively, in their earlier stages in hanging drops. Later, they are quite different. The early conidiophores of the Isaria may lack prophialides, and thus resemble to some extent those of the Cordyceps, though on the latter, the phialides as a rule are not so regularly arranged in whorls. But the conidiophores which occur on the Isaria clava are furnished with whorls of prophialides, each prophialide bearing a cluster of phialides with apical chains of conidia. De Bary's figure, 165 E, as far as regards the branching of the conidiophore, might be matched by the early conidiophores of *I. farinosa* in culture, but the chains of conidia are irregular, and on some of the phialides the conidia are in more or less globose heads, resembling in the latter respect an Acrostalagmus.

In 1894, G. F. Atkinson published a paper, "Artificial cultures of an entomogenous fungus," in *Bot. Gaz.* XIX, 129–45, with three plates. From his figures, he was evidently dealing with *Isaria farinosa*, as understood in this country. He did not obtain any perithecial clavae in culture.

In 1895, R. H. Pettit published Studies in Artificial Cultures of Entomogenous Fungi, Bull. No. 97, Cornell Univ. Agric. Exp. Sta., Bot. and Ento. Divisions. One of the fungi was Cordyceps militaris, cultures of which were made from ascospores (part-spores). The following is

taken from Pettit's account:

"Germination from these swollen spore segments takes place by the production of germ tubes at one or two points. These soon become branched....The threads are strongly segmented and the branches are strongly constricted at the base. In some cases a healthy thread suddenly becomes constricted and produces an aborted apex of less than half the diameter of the ordinary thread. The aborted portion is usually curled....In about four days the growth appears above the surface of the agar. A strong white cottony growth appears, forming a colony circular in form. At the end of about six days the conidia appear. Short sterigmata [i.e. phialides] are borne near the ends of the long cottony threads. They are irregularly arranged either in an opposite or an alternate manner. They are flask-shaped and slender and sometimes forked. The conidia are nearly spherical and are borne in short chains of three or four at the ends of the sterigmata, or at the end of a long thread. The chains are seldom seen, for they almost invariably collapse, leaving the conidia in balls at the ends of the sterigmata."

Pettit also grew the fungus on potato in tubes. The mycelium on the surface of the potato, and the potato itself, were coloured pale orange or brilliant chrome-yellow, wherever they touched the glass. In a half-litre flask of potato, what was probably the beginning of a perithecial clava, deep reddish orange in colour, was observed at the end of about three months. No *Isaria* forms were obtained.

Pettit's figures show short phialides, narrow flask-shaped, scattered along the hyphae, sometimes opposite, sometimes alternate, sometimes in whorls of three. Two phialides may occur on the apex of a short lateral branch of the main hypha (Pettit's forked sterigma), and a phialide may be produced into a long slender thread. The conidia are shown in short chains, or in a small head at the apex of a phialide.

On comparing these figures with those of Tulasne and de Bary, it is seen that they are essentially the same. De Bary's figure, 165 E, shows apparently a more definite conidiophore, but the phialides are similar, in whorls of three, and the conidia are in heads at the apices of the phialides or in irregular chains. The latter are evidently not the definite chains of a *Spicaria*.

Specimens of *Cordyceps militaris* were collected at Austwick, Yorks., in September 1934. Spore prints were obtained the same night, and

when the specimens were left to dry, the ascospores were extruded in long tendrils which formed fleecy masses over and around the clavae. Cultures were made both from the part-spores in the spore

print and from those in the extruded tendrils.

In general, the part-spores germinated readily in hanging drops of water in damp cells. Some of them retained their cylindrical shape. Others assumed an irregular, elongated pentagonal shape, caused by an angular bulging out of one of the longer sides near one end. Some of the part-spores became oval, but these were not observed to germinate.

In the most general method of germination, a short chain of conidia was produced directly from the part-spore, usually from one of the original corners (as seen in profile). Sometimes two chains were produced, sometimes three, from the original corners, or a chain might arise about the middle of the longest side. The first conidium formed was pyriform, with a subacute base, $3 \times 1.5 \mu$, but the succeeding conidia were subglobose, $2 \times 1.5 \mu$.

Other part-spores produced a stout hypha, about twice the length of the part-spore, from one end, like a prolongation of the original spore, and then tapered into a conical phialide, which produced a

chain of conidia at the apex.

In others, a hypha was similarly produced, but ran through the hanging drop for a considerable length, bearing laterally short, scattered, cylindrical phialides, up to 7μ long, each bearing a chain

of conidia at the apex.

Part-spores were also sown on oatmeal agar slants in tubes. Growth was vigorous, and soon the slant bore a thick, greyish white covering, loose and woolly internally, but with an even surface. At the upper edge of the slant, the growth was whiter and somewhat floccose. When old, the covering collapsed into a thin film and became cream-coloured. There was no general growth of erect conidiophores over the surface, nor did it become mealy. The reverse became yellow and then orange, especially where in contact with the glass. No *Isaria* forms have developed in these cultures.

Over the surface of the slant there are no definite conidiophores. Phialides are borne laterally on indefinite hyphae which run in and over the mass of mycelium. These may be alternate, or opposite, or clustered in small groups, sometimes in whorls of three or four, not always at a septum. The phialides are narrow flask-shaped or conical, up to 9μ high, 1.5μ diameter below, tapering to the apex. Sometimes a phialide is prolonged into a very fine hypha, up to 50μ or

more long.

At the upper edge of the slant, the hyphae more closely resemble conidiophores. They may have a terminal phialide, or a terminal cluster of two or three, with scattered phialides, or whorls of phialides below. The whorls, however, are often irregular, consisting of about four phialides arising at one level round the hypha, with one or two a short distance above and below. Sometimes a whorl consists of three phialides and a branch, instead of four phialides, as in de Bary, Fig. 165 E, though the branch may be a simple hypha only. There

are no prophialides.

The conidia are produced terminally on the phialide. After the formation of the first conidium, it is pushed aside by the growth of its successor. The conidia are strongly mucilaginous, and adhere to one another and to the apex of the phialide, so that a globose head, like that of a *Cephalosporium*, is formed. Alternatively, the conidia may adhere to one another in a single or double chain. In some instances, the mass of conidia arising from one phialide takes the form of a globose head, surmounted by a chain; apparently, in these cases the conidia first formed assumed the chain arrangement, but those formed later grouped themselves round the apex of the phialide.

There is no organic connection between the conidia in a chain, as there is in *Spicaria* and *Penicillium*. They are not truly catenulate, but simply adhere by virtue of their mucilaginous coat. That is very evident when the conidia are mainly oval or subpyriform, as they then adhere to one another with their longer sides in contact, *i.e.*

transverse to the direction of production.

The conidia from phialides over the general surface of the slant were chiefly globose, $1.5-2\mu$ diameter, with some oval, up to $3\times2\mu$, and a few subpyriform. At the upper edge of the slant they were chiefly oval, or subpyriform with one end acute, $2.5-3\times1.5-2\mu$.

De Bary noted that the first conidium produced by a phialide was cylindric, like the cylindric conidia found in the body of the insect but shorter. That phenomenon occurs in culture, but apparently not universally. One would not, however, call this first conidium cylindric. It is rather pyriform, with a truncate base, up to 6μ long and 3μ diameter. Frequently this large conidium is perched on the top of a globose cluster of conidia (compare de Bary, Fig. 165 E), or at the apex of a chain of conidia. More peculiar is the fact that in the latter case, when the conidia in the chain are oval and stand transverse to the direction of production, the large terminal conidium stands in the direction of production and consequently at right angles to them.

The results detailed above are in agreement with those obtained by Tulasne, de Bary, and Pettit from cultures of the ascospores of Cordyceps militaris. They also agree with those obtained by Mr E. W. Mason, of the Imperial Mycological Institute, in unpublished research on the same subject. The conidial stage of C. militaris is a Mucedine, with phialides arranged laterally on indefinite hyphae, though towards the ends of the hyphae they may be grouped so as to simulate

a branched conidiophore. The conidia are borne terminally on the phialide and usually adhere in a globose head. The fungus is perhaps best regarded as a *Cephalosporium*. It is certainly not *Spicaria* (*Isaria*) farinosa, which has regular whorls of prophialides and broader flask-

shaped phialides, with truly catenulate conidia.

Tulasne and de Bary were mistaken in identifying the conidial stage of Cordyceps militaris with Isaria farinosa. In other respects their morphological results agree with those of subsequent workers. The only detail which conflicts with this conclusion is de Bary's experiment in which the perithecial clavae of Cordyceps militaris and large Isariae developed from the same pupa. With regard to that there are two possible explanations. The first is that the larva was infected by Isaria farinosa, before de Bary infected it with ascospores of the Cordyceps. The second is that the conidial stage of C. militaris may, under some conditions, assume an Isarioid shape, just as Isaria farinosa may occur as a simple Spicaria or as an Isaria. But as such an Isarioid form has not occurred in culture, nor been found on a pupa in nature, this second explanation would appear improbable.

As noted by Tulasne and de Bary, the conidiophores of Cordyceps militaris may occur on the mycelium on the larva or pupa which bears the perithecial clavae. Apparently they are not always present, or rather, I have failed to find them sometimes. Those I have seen are conical phialides, up to 12μ high, $1-1.5\mu$ diameter at the base, tapering to the apex. They are lateral on the hyphae, or two or three are situated at the apex of a short lateral branch. In one instance, a group of three phialides was seen on the apex of a hypha. The conidia adhere in short chains, or in a small cluster, at the apex of a phialide, and are globose, 1.5μ diameter, or oval, $2 \times 1.5\mu$.

During the last five years, I have made numerous cultures of *Isaria farinosa* with the object of ascertaining whether any specific differences could be established between the different forms of that species, or between the examples which occurred on different hosts. Apart from the fact that *Isaria farinosa* has a definite *Spicaria* conidiophore, cultures of it differ in appearance from those of the conidial stage of *Cordyceps militaris*. They are much looser and more woolly at first, and soon become covered with conidiophores and conidia which give them a mealy appearance. Ultimately, on oatmeal agar or similar media, they produce *Isaria* clavae.

Isaria farinosa has been identified on Lepidopterous larvae and pupae, Hymenoptera, Coleoptera, Aphides, Diptera, and Arachnida. On the larger of these it forms conidial clavae, but on the smaller, and sometimes even on the larger, it produces only a covering of conidiophores. Now that its association with Cordyceps militaris has been terminated, one has less difficulty in accepting the identity of the fungi on these different hosts. It is evidently an omnivorous

entomogenous fungus which can attack insects of all kinds. On Hymenoptera, it has been named *Coremium Swantonii* A.L.Sm.

SUMMARY

The conidial stage of *Cordyceps militaris* is a *Cephalosporium*, which occurs on the mycelium on the larva or pupa bearing the *Cordyceps*. It has not been known to produce an *Isaria* form in nature or in culture.

Isaria farinosa is a fasciculate Spicaria, and may occur as an Isaria or as a simple Spicaria. Its perithecial stage, if any, is unknown. It

has no relation to Cordyceps militaris.

Cordyceps militaris attacks the larvae and pupae of Lepidoptera. There are two records of its occurrence on Coleoptera—on the remains of a cockchafer in the wood of Bailly, Aube, France (Briard), and on a cockchafer, Aude, France (Roumeguère)—but these are generally considered to be erroneous.

Isaria farinosa is a general entomophyte, and is known to occur on Lepidoptera, Hymenoptera, Coleoptera, Diptera, Aphides, and

Arachnida.

Cordyceps militaris of the United States of America is the same as the European species.

SPORES AND SPORE GERMINATION IN WILD AND CULTIVATED MUSHROOMS (PSALLIOTA SPP.)

By DOROTHY M. CAYLEY

(John Innes Horticultural Institution, Merton)

(With Plate III and 2 Text-figures)

 ${
m H}_{
m ITHERTO}$ the classification of the various forms of field and cultivated mushrooms has been based on external morphological characters only, and no account has been taken of the number of spores on the basidium. It has been known for some time that the typical wild species of *Psalliota campestris* and *P. arvensis* have four-spored basidia, with occasional exceptional two-spored individual caps. Sachs in 1868 (see English edition, 1882, Fig. 227) figures two-spored basidia of Agaricus campestris, and Buller (1909) records finding two-spored individuals in the wild on manured ground in the campus of the University of Manitoba, but he states that "they differed considerably from the wild field mushrooms of England, in that they were more scaly, browner and possessed relatively very shallow gills". Buller's specimens were evidently not typical Psalliota campestris, and unfortunately the origin of Sach's specimen is not known. If two-spored forms of typical P. campestris occur in the wild they are rare, and are possibly haploid fruiting bodies of the four-spored type.

It has been stated by Buller and others that the cultivated varieties have two-spored basidia. It is true that two-spored basidia predominate, but, on examination, only one out of three cultivated varieties on the market in this country has shown uniformly two-spored basidia; in the others an appreciable percentage of three-spored basidia occur together with a few one- and four-spored. The variable basidia are not evenly distributed over the gills but are found in patches; Pl. III shows two photomicrographs of the young gills of the fuscous (fig. 5) and the white non-fragrant (fig. 4) cultivated varieties. None of the cultivated varieties examined has shown uni-

formly four-spored basidia as in the wild species.

The three different varieties of cultivated mushrooms examined

and tested for spore germination are as follows:

(1) The fuscous variety on the English market; coarse, thick fleshed, with thick bulbous stipe, tough and tasteless when cooked; basidia and spores variable, basidia one- to four-spored (Pl. III, figs. 1, 1 a).

(2) The white non-fragrant variety, also on the market, softer in texture, flesh thinner than in (1), closely resembling the wild P. campestris; slight but delicate flavour when cooked; basidia and spores variable, basidia one- to five-spored (Pl. III, figs. 2, 2 a).

(3) The white fragrant variety, closely resembling *P. campestris* but rather drier and firmer in texture, with a strong smell when fresh; good texture and flavour when cooked; basidia uniformly two-spored.

In an appendix at the end of this paper full technical descriptions of (1) and (2) are given by Miss E. M. Wakefield. Unfortunately the white fragrant variety (3) has been met with only once; it was bought with stipe cut short, and it has not been possible to procure any more perfect specimens. A full technical description of this and other forms met with under cultivation will therefore have to be deferred to a later publication.

Nothing is definitely known as to the origin of these cultivated forms. The Americans believe that the coarse fuscous variety, which they call *P. brunnescens*, originated in this country and was selected presumably from spawn gathered in the wild; but the wild species *P. brunnescens* is not common either in this country or in the States, and there is no record of its being introduced into cultivation.

There are, however, three records of a form very similar to our fuscous variety occurring in the States. Atkinson (1906) obtained material for his investigation on the development of Agaricus campestris from cultures grown in a greenhouse of a two-spored variety of A. campestris known as "Columbia", sold by the Pure Culture Spawn Co. of Missouri. This variety is figured and closely resembles the fuscous variety described by Miss Wakefield in the appendix. The stipes of mature pilei of "Columbia" as figured by Atkinson are, however, not bulbous. This may only be due to cultural conditions as the immature unexpanded specimens clearly show the typical bulbous stipe.

Atkinson states that he twice found two-spored Agaricus closely resembling certain cultivated forms growing spontaneously in the open; on a lawn which had been mulched with horse manure, and on a hill side of a wooded ravine in the campus of Cornell University.

Murrill (1914) found a number of caps of a fuscous *Psalliota* on an old heap of manure in Bronx Park, New York. He described and figured it in *Mycologia* (1914), and gave it the name of *Agaricus campestre hortensis*. His photographs are not quite clear, and his specimens appear to differ slightly from the common fuscous form on our markets. They are, however, not at all unlike.

F. C. Stewart (1929) purchased spawn from an American Spawn Co., alleged to be the "cream white" variety, but all the pilei produced by this spawn were, as he says, "altogether different from the common mushroom", and, judging from his figures and description,

there is no doubt that he was dealing with the fuscous variety common in this country. Stewart sent specimens to Dr Kauffmann who, after careful examination, expressed the opinion that they might belong to P. brunnescens, a species described by Peck (1929), and that he knew of no other species to which the specimens approached more closely; and further that this fuscous form should not be considered as a variety of P. campestris, P. arvensis or P. subrufescens. Murrill also thought that his form seemed very near P. brunnescens, and states that it is often cultivated but rarely wild. Peck's figures and description, however, differ in several respects from the fuscous form figured and described by Stewart.

There is also nothing definitely known about the origin of the two white cultivated forms, the fragrant and non-fragrant, but they must

have originated at some time from wild species.

The two-spored cultivated varieties are not haploid forms of the four-spored species. The young basidia are binucleate, karyogamy occurs in the basidium, followed by two meiotic divisions resulting in four daughter nuclei, which can be seen distinctly in a resting state just as the sterigmata are about to develop. Sass (1928) describes the passage of two nuclei into each spore in the two-spored form he investigated, and Colson (1935) the same for the two-spored form, and one nucleus into each spore in the four-spored. The nuclei divide again once in the young spore.

SPORES

A number of spores of *Psalliota campestris*, *P. arvensis*, and the several varieties of cultivated mushrooms have been stained and examined for nuclei.

The technique is as follows:

A small platinum loopful of egg albumen, such as is used for mounting microtome sections, is placed on a clean grease-free slide, and spores from a dry spore-trace placed in the drop. The drop is then spread about 1 in. along the slide with a clean glass rod, to ensure the film being thin and even. The slide is then allowed to dry partially for twenty-four hours, protected from dust.

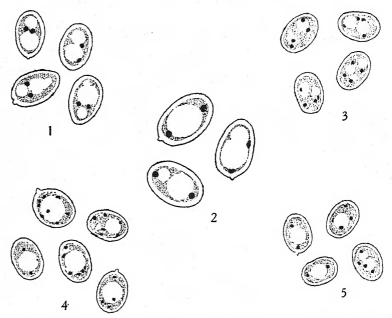
Then the slide is fixed in a fixative recommended by Sass (1929):

1 per cent. glacial acetic acid ... 40 c.c. Commercial formaldehyde... 10 c.c. 95 per cent. alcohol... ... 50 c.c.

for twelve to twenty-four hours, rinsed thoroughly in 50 per cent. alcohol and taken down to water. This fixative hardens the film and the spores do not wash off.

The spore walls of *Psalliota* are dark coloured and somewhat thick, more especially in *P. arvensis*. For the decolorisation of the spore walls

a modification of the method recommended by Brunswik (1924) should be used. Brunswik differentiated with Eau de Javelle after staining with Heidenhain's haematoxylin, but it was found that clearer and better preparations resulted if the process is reversed and the spore walls decolorised before mordanting and staining. The preparations should be left in the Eau de Javelle for half to one hour or longer, until the walls are completely decolorised, then washed thoroughly in water, mordanted in 3 per cent. iron alum for three to four hours, stained overnight in 0.5 per cent. haematoxylin and



Text-fig. 1. 1, wild Psalliota campestris; 2, wild Psalliota arvensis; 3, cultivated white fragrant form; 4, cultivated white non-fragrant form; 5, cultivated fuscous form. ×2150.

differentiated in the usual way in 2 per cent. iron alum, until nothing but the colourless spore outlines can be seen under the low power of the microscope. Differentiation requires a little practice, as the nuclei are not visible under a low magnification.

The results obtained from the examination of these spores confirm Sass's observations (1928) on the two-spored form he worked with, and Colson's results recently published (1935), with two- and four-spored forms of *Psalliota*, namely that spores from two-spored basidia are quadri-nucleate (Text-fig. 1 (3)) and those from four-spored basidia binucleate (Text-fig. 1 (1), (2)).

The spores of the fuscous cultivated form with variable basidia (one- to four-spored), showed both binucleate and quadrinucleate spores in the same spore-trace (Text-fig. 1 (5)). The white non-fragrant cultivated form, also with variable basidia, but mainly two-spored, showed spores with from two to eight nuclei (Text-fig. 1 (4)).

Rough counts made in microscope fields taken at random on young gills of the fuscous and non-fragrant white forms are given

in Table I.

Table I. Number of spores on basidia in cultivated forms

	Basidia.	Percentage					
Pileus	counted	I-spored	2-spored	3-spored	4-spored	5-spored	
		white	non-fragran	t iorm			
Ι	743	4.8	88	6.6	0.2		
\mathbf{II}	359	10	85	5			
Total	1102	6.7	87.2	5 ∙6	ი∙36		
		F	uscous form	ı			
1	250	6.8	37.6	48•4	7.2		
II	250	1.2	49.2	42.8	6.8		
III	950		62.6	29.1	4.4	0.3	
Total	1450	3·4 3·6	56	34.8		0.5	
	-430	3 0	30	34.0	5.3	0.2	

Spore measurements

Psalliota arvensis		•••	•••	$8.9 - 11.1 \times 6.4 - 7 \mu$
P. campestris	•••			$7.6 - 8.9 \times 5.1 - 6.4 \mu$

Cultivated varieties. Basidia two-spored or variable:

Fuscous	•••	$5.7-8.9 \times 5.1-7 \mu$
White non-fragrant		$6 \cdot 4 - 8 \cdot 9 \times 5 \cdot 1 - 7 \mu$
White fragrant (two-spored)	•••	$6\cdot 4-7\cdot 6\times 5\cdot 1-5\cdot 7\mu$

SPORE GERMINATION

An enormous amount of work has been done by various investigators on the conditions conducive to the germination of spores of *Psalliota*. A good review of the literature up to 1924 can be found in Falck's paper (1924) and need not be gone into *in extenso* here.

Hoffmann (1860) is probably the first investigator who succeeded in germinating spores under controlled conditions, in water or damp air. He does not appear to have had any difficulty and remarks that there is nothing unusual about the germination of Agaricus campestris.

Attempts at mushroom culture under controlled conditions were first started in France. Chevreul (1861) and others tried test-tube cultures with very variable and uncertain results.

In America Duggar (1901) and Ferguson (1902) were the first to attempt to germinate the spores of various fungi including A. campestris, on media of known constitution, previously subjecting them to various treatments including artificial digestive fluids. Duggar, in his paper, employed the generic name Agaricus in the sense in which it is usually understood "by those interested in the practical side of the work", and gives no detailed description of the forms or varieties he worked with. This was of course before the days of pure spawn culture.

Both these investigators had very variable results. Ferguson, however, made the important discovery that germination could be stimulated by the introduction of small pieces of vigorous mycelium from a culture of the same species into the spore suspensions. She obtained her spores from sporophores just as the veil was about to break and again when the pileus was fully expanded or nearly so. She observed that spores from one spore-trace from a fully expanded sporophore raised in a conservatory, gave more uniform germination than those from any other spore-trace, but gives no further details as to the origin of the other prints, whether from cultivated forms or from the wild. She also noticed that when a high percentage of germination finally resulted it was always preceded a few days before by the germination of one or more spores. She was wholly unable to account for the irregularities in her results, but suggests that the maturity or other conditions of the spores must be the cause.

In 1924 Falck made a series of very elaborate experiments on the germination of spores of *Psalliota*, treating them with organic and inorganic acids, alkalis, etc., and obtained the best results in a mixture of 0.25 per cent. succinic acid and 10 per cent. malt extract. He repeated Ferguson's experiments and found that germination was stimulated by the introduction of mycelium of the same species. But, again, he gives no definite description of the form he dealt with.

After Falck, little further work appears to have been done on germination until Lambert in 1929 succeeded in isolating single spores of a cultivated variety "snow white", and in obtaining sporophores from single spore cultures, and from all possible combinations of them.

In 1930 Hein was able to germinate spores in distilled water, dung decoction and various synthetic media. The spores were obtained from mushrooms grown for experimental purposes in a specially constructed house. He gives no description of the variety used, but in all probability he was dealing with one or other of the cultivated forms.

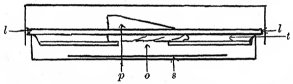
The main object of this paper is to describe a method for spore collection and a simpler and fairly reliable method for inducing germination in the cultivated varieties but more especially in the wild species P. campestris. The spores of the cultivated varieties germinate

more readily than those of the wild species.

Without previous treatment, the spores require an incubation period of from six to seven days before they germinate, even under the best conditions; and, in order to avoid contaminations, the less they are handled before being placed in the solution in which they

are to germinate the better.

Previous preliminary experiments on spore germination in Knop solution only, showed that the condition of the sporophore when the spore-trace is taken is of prime importance. The first shed spores of an immature sporophore do not germinate under artificial conditions. The pileus must be fully expanded and the gills umber, and consecutive traces should be taken from the same pileus at intervals of twelve to twenty-four hours. Therefore, it is advisable when gathering mushrooms in the open, not to gather the cap only, but to cut out the sod on which the cap is growing, so that it can continue to develop fairly normally for some days. The sod can either be left out



Text-fig. 2. Diagram of Petri dish fitted with tin tray and slide, and with sector of pileus in position for collecting spore trace. *l*, lip of tray resting on the edge of lower half of Petri dish; o, opening in the tray; p, sector of pileus; s, slide; t, tray.

of doors in a sheltered place, or brought into the laboratory and placed under a bell jar, which should be slightly tilted to ensure adequate ventilation without risk of drying out. If it is not possible to cut out the sod, the stipe should be pulled up gently and not cut. The pileus can be kept in a fresh condition by placing the base of the stipe on damp cotton wool in a small tube covered with a bell jar, again giving adequate ventilation. Sectors of the pileus can then be cut out when required.

The following is a method of collecting spore-traces free from

contamination:

The quarter pound squat tins of Player's "Country Life" tobacco have lids which fit into a 4-in. Petri dish, so that the lip of the lid rests on the edge of the lower half of the dish, leaving a space of about $\frac{1}{4}$ in. between the lid and the base of the dish. The decoration can be easily removed with methylated spirit and a rectangle cut out of the centre of the lid. A sterile slide is then placed in a sterile Petri dish and the tin tray flamed and placed over the slide. A sector is then cut out of the pileus and placed over the slit in the tray and the lid of the dish replaced (Text-fig. 2). Sectors of large pilei do not

allow of the lid of the dish fitting down closely, but as long as the sector is large enough to cover the slit completely there is no risk of outside contamination of the trace. The slide with the spore-trace can be removed after the required interval and replaced by another, and a fresh sector of the pileus placed on the tray, and so on, until spore discharge ceases.

Before storing, the slides should be allowed to dry for twenty-four hours, then another sterile slide placed over the spore-trace and the two slides fixed together at one end with a sticking label, leaving the other free for opening when required. The slides should be wrapped tightly together in cellophane or clean paper, and stored in a dry

cool place.

The spores of *P. campestris* remain viable for about six months, those of the cultivated varieties rather longer. Spores of the fragrant white variety have germinated after eight months and one spore-trace of the fuscous form gave some germination after fifteen months.

METHOD FOR GERMINATING THE SPORES

A hollow-ground slide is placed in a Petri dish on a circular piece of filter paper with an oblong cut out of the centre to coincide with the hollow in the slide. This eliminates the necessity of removing the slide from the dish for examination under the microscope. The dish is then sterilised. When sterile, a few drops of Knop solution are pipetted into the hollow of the slide, and a loopful of spores from a trace gently dipped into the liquid and not stirred, so that a number of spores remain floating on the surface. Although germination generally begins amongst the submerged spores, the floating spores appear to do better and grow faster.

The filter paper is then moistened with sterile water, the dishes placed on a sheet of glass under a bell jar lined with damp blotting paper and left to incubate undisturbed for about seven days at a temperature of 26–27° C. The Petri dishes should be marked with a wax pencil and not with sticking labels, as *Penicillium* and other impurities develop on the paper in a saturated atmosphere and considerably increase the risk of contamination when the dishes are

opened.

Small pieces of vigorous mycelium may be placed at the edge of the Knop solution either at the time of sowing or seven to ten days after. The latter method was found to be the better in practice.

The objections to inserting the inoculum at sowing are that during the incubation period of seven to ten days the mycelium spreads all over the surface of the Knop solution and may absorb most of it, and also, when spores from a trace are capable of germinating without any inoculum, germination may be delayed for a few days. If the inoculum is introduced after ten days, germination often starts within twenty-four to forty-eight hours and is much more even. The stimulus set up by the inoculum hardly extends beyond the tips of the hyphae, but once a few spores have germinated, a secondary stimulus is set up and germination spreads throughout the liquid.

If the sporelings are required for growing on, they should be taken out of the Knop solution as soon as possible and spread on a solid or semi-solid medium, from which they can be picked off singly or in

groups and transferred to slopes of oatmeal agar.

Spore Germination tests

Table II gives two series of germination tests with spores of the wild species *Psalliota campestris*, in Knop solution, with and without

inocula from stipe cultures of the same species.

LS 7, LS 7a, and LS 7b are three consecutive spore-traces from a mature pileus with umber gills, taken at the intervals stated. It will be seen that in Knop solution only, without any inoculum, the spores of the first spore trace (LS 7) gave the best results on the eleventh day in series A, and some germination on the eighth day in series B. The third trace (LS 7b) gave little or no germination in B until the seventeenth day.

With the addition of inoculum at sowing, the second trace proved the best on the tenth day in series A, and gave some germination on the eighth day in series B. With the addition of inoculum on the tenth day the third trace (LS 7b) gave good germination in twenty-four hours in series A, and in B a percentage equal to that of LS 7

in series A.

The Knop solution in both series was not freshly made, but had

been kept in tubes for some weeks before use.

It must be pointed out that these percentages were obtained from counts in microscope fields taken at random in the region within the influence of the stimulus set up by the inoculum, either underneath or round the edge of the mycelium. They are consequently merely rough estimations and only serve as a means of comparison between the effects produced by the different treatments. A true estimate of the percentages would entail spreading the spores from the suspensions daily on a solid or semi-solid medium for several days in succession, allowing them to incubate and counting them. Neither time nor incubator space would allow of this method being adopted with a large number of spore-traces; but, in one single instance, the percentage of germination was considerably increased by plating and counting the spores some days after germination had started in the suspensions.

Psalliota does not do well in a solution. After the germ tubes have

Table II A. Germination tests in Knop solution.

	Å	Inoculum Inoculum added at added after sowing 10 days	A little	+ In patches	#
	14th day		# 46.8 %	++	+ +
		No. inoculum	+	A little in patches	1
p_{lq}	y	Inoculum added after No added at added after 10 days inoculum sowing 10 days	Medium Knop. Occasional spore	Occasional spore	48.4 %
es 3 days e	rith day	Inoculum added at sowing	s moculum. +	++ 26 %	++ 20.6%
Spores 7 weeks 3 days old		No inoculum	n F. campestr. # 45.7 %	1	I
S	a 6	Inoculum Inoculum added at added at sowing 10 days	r. campesuris Wil	1	Occasional spore
	10th day	Inoculum added at sowing) + ₁ 6	++	+
	-	No. inoculum	1	1	1
	Date	Spore-trace, P. campestris	LS 7, gills umber 1st 16 hr.	LS 7a, 16-40½ hr.	LS 7b, 40½-64 hr.

Table II B. Spore germination tests in Knop solution.

	17th day	Inoculum added at sowing	25 %	50 %	% 5.65
		Inoculum added after 10 days	56 %	34.2 %	27 %
	day	Inoculum added at sowing Knop.	1	Nothing further	1
Spores 8 weeks 6 days old	roth	No inoculum ulum. Medium	Occasional spore+ inoculum added	I	Inoculum added
	8th day	Inoculum added at sowing P. campestris inoc	+	A little+ in patches	1
		No inoculum . campestris with.	+ Occasional spore	1	1
	7th day	Inoculum added at sowing P	+ Occasional spore	1*	I
	711	No inoculum	1	1.	1
	Date	Spore-trace	LS 7	LS 7a	LS 7 <i>b</i>

attained a certain length, the spores are either pulled up to the surface of the liquid and there get caught up in the inoculum, or, if they remain submerged, the germ tubes die and disappear. If the inoculum is very vigorous, it is often necessary to draw it aside to expose the spores lying underneath, and a certain number of sporelings are drawn away with it. This mainly accounts for the decrease in the percentages in series B after the twelfth day. Also after the germ tubes of the floating spores have attained a considerable length they become so entangled that they cannot be counted.

The patchy germination recorded in this table is most probably due to a secondary stimulus set up by the scattered germinating spores seen on the eleventh day and not to the stimulus set up by

the inoculum.

Table III gives a more comprehensive series of germination tests, with inocula of the same and different species. The same spore-traces LS 7, LS 7a, LS 7b as in Table II are included, when the spores were four months old; also two spore-traces from another pileus of *P. campestris* S 32, S 32a, gathered when the gills were not fully ripe but pinkish umber in colour. CP 4a is the second spore-trace of the white fragrant cultivated variety.

Table III. Germination tests with inocula of the same species and different species

			· op corec		
			species apestris	Different species <i>P. arvensis</i> cultivated	
Spore-trace P. campestris	No inoculum	Inoculum same species added at sowing	added after 7 days	Inoculum P. artensis after 7 days	Inoculum fuscous form after 7 days
		Date: 10th day after sowing			
		Fre	sh Knop 23. ii	• 35	
LS 7, + 1st 16 hr. gills umber	+	+	A few spores	+ Occasional spore	36·3 %
LS $7a$, $+ 16-40\frac{1}{2}$ hr	Occasional spore	- -	+	+ Occasional spore	32·9 %
LS $7b$, $+40\frac{1}{2}$ -60 hr	+	_ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~	+	+ Occasional spore	+ Occasional spore
S 32, 1st 16 hr. gills pinkish umber	+ 1 spore	+ 1 spore	. -	+ Occasional spore	+ Crystals; a little germ.
S 32 a, 16–29 hr.	-	+ A few spores	+ A little germ.	<u> </u>	+ Crystals; fair germ.
		White fra	grant cultivat	ed variety	
CP 4a, 17-37½ hr. gills umber	Contaminated with bacteria	l +	# 43 %	A few spores	+

Table III (continued)

			species	Different species P. arvensis cultivated	
Spore-trace P. campestris	No inoculum	added at sowing	Inoculum same species added after 7 days	Inoculum P. arvensis after 7 days	Inoculum fuscous form after 7 days
			12th day after sh Knop 23. ii		
LS 7, + 1st 16 hr. gills umber	+	+	+	+	′ 50•8 %
LS $7a$, + $16-40\frac{1}{2}$ hr	Occasional spore	_	+	Occasional spore	# 52·8 %
LS 7 b , $+40\frac{1}{2}$ -60 hr	. +	+ Occasional spore	+	+ Occasional spore	28 %
S 32, 1st 16 hr. gills pinkish umber	+	Nothing further	+	Occasional spore	+ Crystals; nothing further
S 32 a, 16-29 hr.	+	+ Occasional spore	+	_	+ Crystals; nothing further
		White fra	grant cultivate	ed variety	idithei
$CP 4a$, 17-37 $\frac{1}{2}$ hr. gills umber	Bacteria	Crystals; nothing further	43·I %	Early # stages 44.9 %	# 40·0 %
		Date: 1	6th day after		
		Fres	sh Knop 23. ii	· 35	
LS 7,+1st 16 hr. gills umber	+ Early stages 17.7 %	# Early stages 30.2 %	+ Nothing further low %	+ 11·2 %	48.8 %
LS $7a$, + $16-40\frac{1}{2}$ hr.	+	+	+ Not countable	+ Nothing further	# 35 %
LS $7b, +40\frac{1}{2}$ -60 hr.	Low %	+ Nothing further	21.9 %	26 %	28.5 %
S 32, 1st 16 hr. gills pinkish umber	+ Nothing further	+ Nothing further	+ Early stages	+ Nothing further	+ Crystals; nothing further
S 32a, 16–29 hr.	+ Nothing further	+ Nothing further	+ Nothing further	+	+ Not countable
	1	White frag	grant cultivate	d variety	
CP 4a, 17-37½ hr. gills umber	Occasional spore; bacteria	+ Nothing further	#Floating spores 33.4 %	36·8 %	# 41 %

LS 7, LS 7a, LS 7b, spores 4 months old. S 32, S 32 a, spores 4 months 10 days old. CP 4a, spores 8 months 18 days old.

It will be seen that all the three traces of LS 7, although older than in Table II, have shown some germination on the tenth day in Knop solution only, but germination has been retarded in the spore suspensions of LS 7a and LS 7b by the inoculum introduced at sowing, and this retardation continues until the sixteenth day.

The inocula of *P. arvensis* introduced on the seventh day do not appear to produce any marked stimulatory effect, and the percentage recorded on the sixteenth day may be attributed mainly to the secondary stimulus set up by the presence of some germinating

spores as in the controls without inoculum.

The stimulatory effect of inocula of the fuscous cultivated form, on the other hand, is very marked on the third day after the introduction of the inoculum, in the first two spore-traces of LS 7, and still more marked on the fifth day. The third trace LS 7b does not respond so readily, and the effect is not more pronounced than with inocula of P. campestris itself. In this series the first two traces have given the best results.

With regard to S 32 and S 32a the spores from pinkish umber gills, the percentages of germination are low throughout. The crystals recorded in the suspensions with the fuscous inocula require some explanation, as they produce an inhibiting effect which has to be taken into account. The aerial mycelium of both wild and cultivated forms of *Psalliota* is thickly encrusted with crystals, analysed by Hein (1930) and found to be calcium oxalate. The fuscous form produces a considerable amount of fluffy aerial mycelium on oatmeal agar. Some of this aerial mycelium was introduced in the inoculum, with the result that crystals were deposited in the Knop solution. These crystals dissolve slowly and completely alter the chemical constitution of the Knop solution and further germination is inhibited.

The effect produced by the different inocula on the fragrant white cultivated form CP 4a is interesting. The fuscous inoculum is not more potent than the inoculum of CP 4 itself, although they are both cultivated forms. Again, the suspensions containing crystals show inhibition of growth and further germination, and accidental contamination with bacteria is also deleterious. In other germination tests with CP 4a, this trace has shown that the spores are capable of

germinating in Knop solution only.

Besides the tests recorded in Tables II and III, many other tests with spores of *P. campestris*, with and without inocula, have shown that success mainly depends upon the maturity of the sporophore when the traces are taken. During spore discharge from one and the same pileus, there appears to be an optimum germination period, either without inocula, or with certain varieties of inocula, and that this period can be shifted by subjecting the spores to other stimuli. Hence the advisability of taking consecutive spore-traces.

During this investigation most of the spore traces obtained by the method described on p. 231 have been free from contamination. Provided fresh, clean pilei are selected and due care exercised, the risk of contamination of spore traces from fully expanded pilei is not great.

Attempts at germinating spores of *P. arvensis* have failed. This is probably due to the lack of spores from mature sporophores, as the spores obtained at the beginning of this investigation were all from immature pilei, and the following autumn no further material was

found.

DISCUSSION

In view of the uncertainty about the origin of the cultivated varieties dealt with in this paper, no object is to be gained, at this juncture, by classing them as varieties of any one or other of the wild species *Psalliota campestris*, *P. arvensis*, *P. brunnescens*, etc., until proof is forthcoming whether they are or are not mere varieties. They are therefore described merely as cultivated forms of *Psalliota*. With the exception of the fuscous form, the other white cultivated forms more nearly approach *P. campestris*, than any other wild species met with so far.

It has been pointed out above that, apart from morphological characters, the main difference between the cultivated forms and the wild species is the number of spores on the basidium, and the number of nuclei in the mature spores. Lambert has shown that the spores of the cultivated "cream white" variety, with two-spored basidia, is monoecious; and since each spore receives two basidial nuclei, the spores are presumably in the diplophase. So far, there is no record of heterothallism in the genus Psalliota, but the fact that, in the four-spored wild species the spores receive only one instead of two basidial nuclei, and must consequently be definitely haploid, suggests that P. campestris and P. arvensis may be haplo-dioecious. Another possible difference between the wild and cultivated mushrooms may be that the cultivated forms can complete their life history from spore to spore as saprophytes, whereas there is some experimental evidence, not yet published, that a form of symbiosis exists between various grasses and the field mushroom P. campestris. Artificial inoculations under sterile conditions have shown that the mycelium can penetrate and live in the roots of grasses for a time without materially affecting the growth of the grass plants. It is not known how long this symbiotic balance persists; but, judging from the examination of roots from sods carrying pilei gathered in the open, the infected roots eventually die and are replaced by new growth in the root system. In the open, however, the grass evidently derives some benefit from the presence of the spawn of P. campestris.

as can be seen on fairy rings formed by this species. The grass on the rings is deeper green and of a more luxuriant growth than either inside or outside the ring. The mycelium of *P. arvensis* will also penetrate and live in grass roots, but in the open, the spawn is so profuse that it sets up physical and possibly chemical conditions, such as impermeability to water and excessive deposits of calcium oxalate, which are detrimental to the deeper rooted grasses.

SUMMARY

The basidia of the two wild species *Psalliota campestris* and *P. arvensis* and three different forms of cultivated mushrooms vary in the number of spores on the basidium. The wild species have four-spored basidia and the cultivated either two-spored or from one- to four-spored.

Only one out of these three cultivated varieties has shown uni-

formly two-spored basidia.

The number of nuclei in the mature spores of the wild species is constant; they contain two nuclei. In the cultivated variable varieties the number of nuclei may range from two to eight.

Methods are given for obtaining consecutive spore-traces from single pilei, for staining spores for nuclei, and for spore germination.

The stage of maturity of the sporophore when the spore-traces are taken is all important.

In conclusion I wish to acknowledge my indebtedness to Miss E. M. Wakefield for her help and for the technical descriptions at the end of this paper; to Mr H. G. Osterstock for the photomicrographs; to the laboratory assistant A. F. Emarton for the photographs reproduced in Pl. III; and to the foreman Mr J. Newell for specimens of the fuscous and non-fragrant white cultivated varieties.

APPENDIX

DESCRIPTION OF TWO FORMS OF CULTIVATED MUSHROOM

By E. M. WAKEFIELD

White form

Pileus up to 7 cm. in diameter, at first hemispherical, then flattened-convex, margin incurved and extending beyond the gills.

Surface of pileus smooth, soft to the touch like a kid glove, under the lens appearing made up of silky fibrils, pure white, but becoming stained with brown when handled. Flesh 12-13 mm. thick in the centre, attenuated towards the margin, white, solid, but less firm than in the brown form, becoming slightly pinkish when cut.

Lamellae crowded, reaching the stem but free from it, rounded behind, narrow, thin, at first pale, clear flesh-pink, becoming purplish

brown (warm sepia (R)) as the spores mature.

Stipe 6-8 cm. long by $1\frac{1}{2}$ -2 cm. broad, white, tinged rosy purple at the apex, smooth with a satiny sheen, equal or slightly incrassated at the base, becoming rufescent when rubbed.

Veil white, membranous, silky, thin and very frail, when ruptured

leaving an irregular annulus on the stem.

Annulus median, white, at first broad and flaring upwards, but

soon collapsing and eventually almost disappearing.

Basidia two-spored. Spores broadly elliptical, very variable in size, and also variable in colour, some being quite pale while others are more or less deep reddish brown.

Spore print dark brown (sepia (R)).

Spores $6.4-8.9 \times 5.1-7 \mu$.

Brown form

Pileus up to 9 cm. broad, convex (flattened and hemispherical) when young, with margin strongly inrolled, becoming convex-

expanded with age, with a slight depression in the centre.

Surface of pileus in the unopened state somewhat tomentose, avellaneous (R) in colour with sometimes, especially towards the disk, slightly broken up into scales. As the pileus expands the scales become more evident, and especially on handling or on drying the colour becomes deeper brown and the scales appear fibrillose and closer adpressed, especially towards the margin. Margin whitish and exceeding the gills, in expanded specimen often recurved, striate rim consisting of fragments of the veil.

Flesh very thick, compact, firm, whitish or tinged pinkish, be-

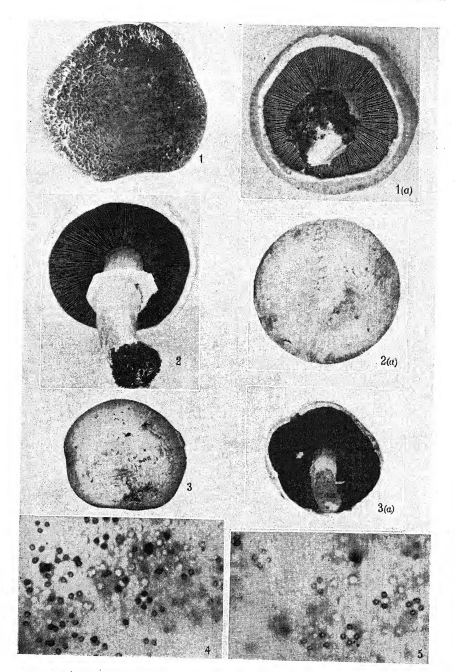
coming brownish when cut.

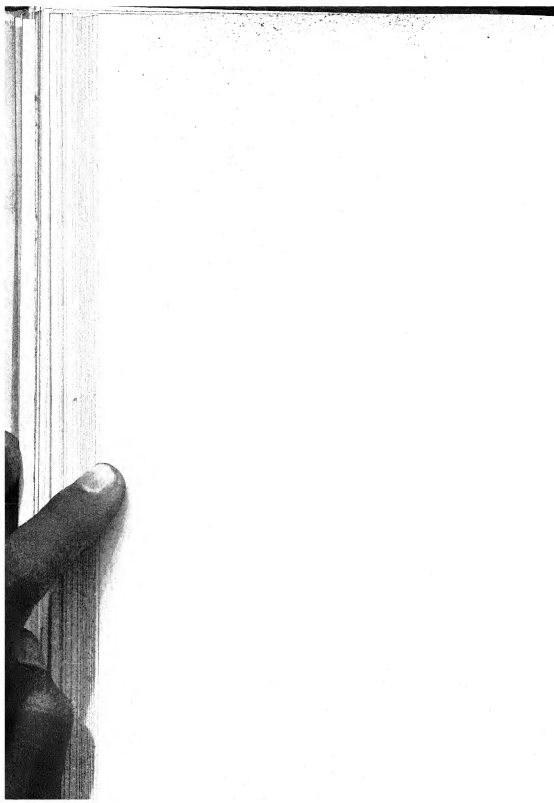
Lamellae crowded, free but reaching the stem, often appearing adnate at first, dull, dirty pinkish brown (approximately vinaceous fawn (R)) with a white edge due to tufts of clavate, hyaline hairs, later deep brown, almost black, narrow compared with the thickness of the flesh.

Stipe usually short and thick, 3-4 (-6?) $\times 2-3.5$ cm. thick, whitish, smooth swollen, sometimes abruptly so at the base, which in indoor specimens may be woolly, solid, or rarely with a slight hollow, becoming brownish when cut or rubbed.

Veil thick, forming a soft, swollen median annulus which appears more or less triangular in vertical section, white and sulcate on the

upper side, sometimes tinged with fawn colour below.





Basidia usually two-spored. Spores broadly elliptical, guttulate, rather thick walled, reddish brown by transmitted light, but giving an umber-brown spore print (sepia (R)).

Spores $5.7-8.9 \times 5.1-7 \mu$.

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EXPLANATION OF PLATE III

Fig. 1, 1 a. Fuscous cultivated variety.
Fig. 2, 2 a. White non-fragrant cultivated variety.
Fig. 3, 3 a. Wild species, Psalliota campestris.
Fig. 4. Photomicrograph of young gill of white non-fragrant variety showing one-to-four-spored basidia. × 140 approx. Fig. 5. Photomicrograph of young gill of fuscous variety showing two- to four-spored

basidia. \times 140 approx.

THE PARASITISM OF MYROTHECIUM RORIDUM TODE

By N. C. PRESTON, B.Sc.

(With Plates IV and V)

During the summer of 1932 it was noticed that certain pansies in my garden appeared to be slowly dying out. First one shoot and then another gradually withered from the base upwards until by about August the whole plant was almost completely dead.

The stems had rotted at ground level and came away readily from the root which was more or less in a sound condition. On the basal portion of the dead shoots thus detached were numerous small black dots surrounded by a distinct white rim which proved to be the fructifications, sporodochia, of a *Myrothecium* closely approximating

to the species usually determined as M. roridum Tode. The constancy with which these fructifications appeared on the dead and dying plants aroused suspicion that the fungus giving rise to them, though generally regarded as a saprophyte, might actually be the cause of the disease. A preliminary test was therefore made, before actually isolating the fungus, by transferring some of the spores from the centre of a sporodochium to the surface of the leaves of a growing potted plant of Viola cornuta. Within three days each of the leaves thus inoculated showed definite lesions in the form of dark purple spots, the centre of which rapidly became dry and brownish. In view of these immediately positive results the fungus was isolated in pure culture.

Pure cultures of the fungus were readily obtained by stirring up spores, taken with a sterile needle from the centre of a sporodochium, in a small quantity of sterile water and spreading a drop of this dilution upon the surface of a previously poured agar plate. After forty-eight hours' incubation at 22° C. the isolated growths could be readily picked out and transferred to agar slopes. In even the initial platings there was extremely little contamination though one or two bacterial colonies appeared. These were also isolated and dealt with separately.

The fungus cultures thus isolated were further tested for any bacterial contamination which might possibly have still been overlooked by taking spores from seven-day growths on malt agar and corn-meal agar respectively, suspending these in sterile distilled water and making poured plate cultures from them with plain nutrient agar.

¹ For this identification the author is indebted to Miss E. M. Wakefield.

These plates were kept at 22° C. and examined frequently. Both plates remained free from any visible bacterial colonies for four weeks, by which time there was a very abundant growth of the fungus mycelium upon them.

BACTERIAL INOCULATIONS

As far as could be readily ascertained the few bacterial colonies referred to above appeared to be those of a single organism and, though such colonies were actually very scarce in the initial plates, it seemed just possible that this organism might have been responsible for the lesions produced in the first rough experiment, and a test was therefore arranged as follows. Three shoots of a healthy plant of Viola growing in a pot were inoculated with the bacterium, while four others were similarly inoculated with spores from a pure culture of the fungus Myrothecium roridum. The inoculum consisted of a suspension in sterile water of spores or bacteria respectively, each being taken from an eight-day-old culture on cherry agar. A drop of the inoculum was placed upon the surface of the plant by means of a sterile platinum loop without any pricking or wounding of the tissues. Each shoot was inoculated upon two isolated internodes. All the shoots thus inoculated with the fungus spores developed characteristic lesions within eight days, while none appeared upon any of those inoculated with the bacterial suspension or elsewhere upon the plant.

INOCULATION EXPERIMENTS WITH THE FUNGUS

The second series of inoculations was carried out on September 9, again on a growing Viola. Two forms of inoculum were used, one a suspension of spores, the other a suspension of mycelial growth broken up in sterile distilled water. The mycelium for this purpose was taken from the fringe of an actively growing culture. Five stems were inoculated with the spore suspension and three stems with the mycelial suspension, the liquid being applied to the plant surface without any scraping or wounding, the plant being afterwards covered with a bell-jar to maintain a moist atmosphere. At the end of three days all the stems inoculated with the spore suspension showed characteristic purple-brown streaks on the internodes inoculated. Of the three stems inoculated with the mycelial suspension one showed a definite lesion comparable with those on the other five, on the second the lesion was apparent but much slighter, while on the third no lesion was visible. All the remaining uninoculated six stems of this same plant, which could be regarded as controls, remained perfectly normal.

This experiment has been described in detail because, in connection with it, a further test was made to eliminate any possibility of

bacterial contamination being a contributory cause of the lesions produced. This was done by placing a small quantity of each of the suspensions actually used for the inoculation in a sterile Petri dish and adding nutrient agar, previously cooled to 40° C. The plates thus prepared remained quite free from bacterial growth. In view of this result and the fact that the bacteria initially occurring naturally in the sporodochia failed to produce infection, it was considered unnecessary to subject the isolated fungus to any further tests of this nature.

The foregoing experiments were followed up by numerous other similar inoculations both on plants in pots and on detached leafy shoots. The stems, leaves, and basal parts were all subjected to inoculation with positive results. The varieties of *Viola* used were a mauve form, "Maggy Mott", a deep yellow, "Chantryland", and the small blue *V. cornuta*. No marked difference in susceptibility between these varieties was recorded.

It is not proposed to detail all the individual experiments, since the majority were carried out in a manner essentially similar to those already described, but reference to any significant diversions from

this method will be made as is necessary.

As regards controls, in the earlier experiments a due proportion of leaves or internodes on the plant used for inoculation were treated with loopsful of sterile water in place of the spore suspension. Later, however, it was deemed satisfactory to regard all uninoculated parts as controls, the inoculated internodes or leaves being carefully marked by loops of thread. Where stem inoculations were made adjacent internodes were not selected for inoculation, the one or more internodes intervening between the inoculated ones thus serving as particularly useful controls. Such control parts invariably remained normal.

STEM AND LEAF INOCULATION OF GROWING PLANTS

(a) Spore suspension applied without wounding or pricking of tissue

Sixteen plants of varieties "Maggy Mott" and "Chantryland" were used in these experiments and ninety-one separate inoculations were made. The results may be summarised as follows:

Internodes inoculated 34; positive infections 25 = 73.5 per cent. Leaves inoculated 57; positive infections 39 = 68.4 per cent. Control parts, as described above, all normal.

(b) Tissues pricked before inoculation. Exp. 14 C

In a single plant, variety Chantryland, the stem and leaf tissues were lightly pricked with a sterile needle and the spore suspension applied in the usual way to the punctured surface.

Five internodes and four leaves (upper surface) were thus inoculated while the same number of each, similarly pricked but treated with drops of sterile water instead of sporesuspension, served as controls.

All the inoculated internodes and leaves showed very pronounced lesions within eight days, the discolouring first becoming visible two days after inoculation. The pricked but uninoculated controls showed no discoloration whatever and remained perfectly normal throughout.

It is of interest to note, in connection with this experiment, that the plant described was one of a series of three, the other two of which were inoculated, unsuccessfully, in the usual way, *i.e.* without pricking. The inoculum for the three plants was prepared from an eleven weeks old culture. The effect on the three individual plants is here shown for comparison:

- Plant A. 3 internodes inoculated without pricking: result 1 positive, 2 negative.
- Plant B. 5 internodes and 4 leaves inoculated without pricking: result all negative.
- Plant C. 5 internodes and 4 leaves inoculated after pricking: result all definitely positive.

The fact that all but one of the A and B inoculations proved negative may perhaps be accounted for by the age of the culture, since in a previous experiment, two days before, it proved similarly inactive giving only two positive results out of ten inoculations.

STEM AND LEAF INOCULATION OF DETACHED SHOOTS

The inoculation of detached shoots of *Viola cornuta* was resorted to at times for convenience.

Only thoroughly healthy strong growing shoots were selected. These were well washed and placed in large covered glass dishes 20 cm. in diameter. The control shoots were sometimes contained in the same dish as those inoculated, sometimes in separate dishes, and in all the experiments here discussed they remained perfectly fresh and normal to the end. Fifteen shoots were used for inoculation.

Internodes inoculated 34; positive 21 = 61.8 per cent. Leaves inoculated 35; positive 32 = 91.4 per cent. Shoots kept as controls 11; all parts remained normal.

In one of these experiments the effect of a very dilute spore suspension was compared with that of a heavy suspension with the following result:

Dish a. 5 shoots. 15 internodes inoculated with weak suspension: positive 3, after 10 days.

10 leaves inoculated with weak suspension: positive 10, after 10 days.

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Dish b. 5 shoots. 11 internodes inoculated (heavy suspension): positive 11, after 10 days.

10 leaves inoculated (heavy suspension): positive 10, after 10 days.

Dish c. 4 shoots. Uninoculated controls. All parts quite fresh and healthy after 10 days.

In both (a) and (b) the characteristic discoloration began to be apparent after three days. In (b), however, the lesions developed much more rapidly; they were very marked on eight out of the eleven internodes within three days, and all were obviously heavily infected within seven days.

INOCULATION OF BASAL PARTS

Since this fungus first came under observation as the cause of a basal rot it was obviously desirable to see whether a rot of this type could be caused by surface inoculation with the pure culture. Some experiments to determine this were therefore carried out, and these will be best described individually.

Exp. No. 5. Two healthy rooted cuttings A and B of Viola var. "Maggie Mott" in pots, were inoculated on February 23, 1933, with material taken from a sixteen days old culture on potato-glycerin agar. A heavy spore suspension was poured down the stem bases of each plant and allowed to run into the surrounding soil and, in addition, some mycelium from the same culture was carefully incorporated with the surface soil in each pot. A third plant C was similarly treated with distilled water to serve as a control, and the three plants were kept together in a cool greenhouse.

On March 27 all three plants, which superficially appeared equally healthy, were lifted and the soil carefully washed from the roots; their appearance was as follows:

Plant A. Main stem with pronounced dark lesion extending from soil level to a height of 2 cm. Over the lower half of this lesion the tissues appeared dark brown, wrinkled and rotting, the upper half of the lesion being a purple-black discoloration. One of the lateral shoots arising 3 cm. above the soil level was similarly blackened over the length of the first internode. The roots and upper aerial parts appeared normal.

Plant B. Apart from slight discoloration the main central shoot appeared sound. Two lateral shoots were characteristically blackened from soil level to a distance of about 2 cm. upwards. Both shoots were rotting at the base. Roots and upper aerial parts were healthy.

Plant C. All parts above and below ground appeared quite healthy. No discoloration apparent.

After examination each plant was transferred to a separate large covered glass dish. When the plants were examined two days later numerous sporodochia of *Myrothecium* were found to have developed upon the basal parts of the stems of plants A and B. Plant C still remained perfectly clean and normal in appearance.

Since the "Maggie Mott" plants were cuttings it seemed possible that they might be more readily susceptible to infection through the callused area than if they had been grown from seed. In further experiments therefore seedlings were used in place of cuttings.

Exp. No. 11. Six healthy seedlings of the yellow Viola var. "Chantry-land", growing in 4-in. pots, were inoculated on August 4, 1934, with a spore suspension of an eight weeks old culture by allowing 3 c.c. of the suspension to run down the lower parts of the stems into the soil. Five similar control plants were treated in the same way with distilled water. The pots were then sunk in soil contained in two large boxes, the inoculated plants occupying one box the controls being in the other, and all were kept in the open.

On August 21 one of the inoculated plants and one control were lifted, washed and examined. Both appeared perfectly healthy and sound at the base. Ten days later (August 31) three more of the inoculated plants were lifted; these also appeared normal, as also did the still growing controls. After examination, these three plants were immediately transferred to covered glass dishes and used for a further

experiment described later.

The remaining two inoculated and four control plants were allowed to grow for another seven weeks. When they were examined on October 18 the stem bases and upper parts of the roots of both the inoculated plants showed distinct signs of rotting from the exterior inwards. In addition to extensive discoloration of the stem bases the main tap root, for about 1 cm. from soil level downwards, was brown and discoloured, the discoloration merging gradually into the white firm tissue below.

All the four control plants were quite normal. Their stem bases were healthy, and it was particularly noticeable that the upper part of the root, which in the inoculated plants was badly affected, was

here perfectly sound and white.

Exp. No. 11 a. The three plants from the previous experiment which had been transferred to glass dishes ten days after inoculation were used. Two of these, in dishes A and B, were further inoculated by running a few drops of fresh spore suspension on to the region between root and stem by means of a pipette, and a small fragment of mycelium was also applied at this point. The plant in the third dish C received no such inoculations and served as a control. On September 10 plant A showed only a scarcely perceptible darkening of the tissues at the point of inoculation, plant B was definitely rotting

at this region, while plant C, the control, remained white and normal.

Exp. No. 12. A single healthy "Chantryland" seedling was inoculated by pouring a spore suspension over the basal part as in previous experiments. A second plant was kept as an uninoculated control. Both plants were washed and examined three weeks after inoculation

when their appearances were as follows:

Inoculated plant. Stem base blackening over a distance of 1 cm. from soil level upwards, the discoloration extending into one of the two secondary shoots. The lesion also continued downwards to a point $\frac{1}{2}$ cm. below the soil level where the discoloration gradually diffused into the sound white tissue of the main root. Sporodochia were present in the middle of the main lesion.

Control plant. No lesion or discoloration was present, the normal greenish tinge of the stem base fading into the clean white of the

healthy root.

Exp. No. 13. Three healthy "Chantryland" seedlings in 7 in. pots were used. From each of the three plants the soil was washed away at one side by means of a jet of water so as to expose the stem base and adjoining roots. One of the plants, A, was then pricked lightly on the exposed surface with a sterile needle and inoculated with spores from an eleven weeks old transfer by means of a platinum loop. The second plant, B, was similarly inoculated but without pricking, the third, C, was pricked but uninoculated. All three plants were washed and examined eighteen days later.

Plant A showed brown discoloration down the stem and root over 1.5 cm. Tissues below ground were clearly splitting and rotting.

Plant B showed a very slight discoloration at the inoculated region but no signs of rotting.

Plant C. Quite normal.

RECOVERY OF THE FUNGUS FROM ARTIFICIALLY INOCULATED PLANTS

In order to establish the parasitic nature of any fungus conclusively it is necessary to be able to reisolate it from the plant lesions produced by inoculation with the pure culture. This was done successfully with Myrothecium roridum. The first plant selected for this purpose was a "Maggie Mott" Viola which had been inoculated on August 27, 1932, and in which the characteristic lesions had been produced upon the stem. Eight days after inoculation a small fragment was removed from the drying centre of each of two stem lesions with the point of a flamed scalpel and the fragments of tissue transferred to two cherry agar plates. A pure growth of Myrothecium roridum, without any trace of contamination, developed upon each of the plates thus inoculated.

The somewhat rough and ready technique adopted in the foregoing experiment did not, however, absolutely preclude the possibility of the transference of a merely surface growth, or even perhaps of the spores of the fungus, to the agar plates, and a more elaborate

procedure was adopted in the following later experiment.

On August 31, 1934, a "Chantryland" seedling was inoculated and kept under a bell-jar until September 4, when definite lesions had appeared on the stems and leaves. One of the stem lesions extended practically the length of an internode, appearing as an elliptical spot, brown and shrunken at the centre and sharply delimited by a purple-black border from the normal green tissue beyond. This internode measured 17 × 3 mm., the actual dimensions of the lesion being 12 × 2 mm. The following day a portion of the stem including the infected internode was cut away, immersed for one minute in 0·1 per cent. mercuric chloride, and thoroughly washed in sterile water. Fragments from the infected spot were then removed with a sterile scalpel and plated on cherry agar.

Two infected leaves, one from this same plant and the other from a second plant inoculated at the same time, were also taken. These were similarly treated with 0·1 per cent. mercuric chloride, washed in sterile water, and portions of the diseased tissue transferred to

agar plates.

From each of the stem and leaf fragments thus plated a luxuriant mycelial growth was obtained and the typical spore masses of *M. roridum* developed after three days.

Description of the fungus

Beyond the bare descriptions of Myrothecium roridum which are to be found in systematic works such as Rabenhorst's Kryptogamen-Flora, I have not been able to trace any literature dealing with this fungus, and no previous suggestion of its being actually parasitic seems to have been made. The following brief description may perhaps, there-

fore, not be out of place.

 \dot{M} . roridum is a member of the Tuberculariaceae, the genus being characterised by the presence of a ring of pure white setae around the margin of the sporodochium. The hyphae, in culture, are hyaline to pale brownish. The spores are straight with rounded ends, continuous and often contain two or sometimes three droplets. They are greenish or of a pale olive tint, appearing jet-black in the mass, and measure approximately $7-8\times 2~\mu$.

The fungus grows well at room temperature on many kinds of artificial media, producing a flocculent growth of pure white aerial mycelium. Sporodochia are usually produced in abundance either on the substratum itself or among the loose aerial hyphae. When

lying separate from one another they appear as jet-black dots surrounded by a white rim. Often, however, they coalesce into larger masses and may sometimes form an almost continuous black line

around the margin of the culture.

The lesions produced by inoculation of leaves or stems of *Viola* appear first of all as dark purple-black spots or streaks which gradually increase in extent. As the lesion advances the tissues at the centre become dry, shrivelled and brown, the outer margin being sharply delimited from the normal green of the leaf by a deep purple-black band.

CONCLUSIONS

From the experiments described it is clear that Myrothecium roridum can function as an active parasite. It is evidently able to enter the unwounded tissues of healthy plants of Viola and eventually to destroy them. Considering the ease with which infection can be secured and the rapidity with which the lesions develop under suitable conditions, it seems evident that this fungus must be reckoned as one of the possible causes of the dying out of violas under cultivation. This view is supported by the fact that the fungus occurs commonly upon dead or dying Viola stems from which it was actually isolated at the beginning of this investigation.

The method of inoculation adopted in the majority of the experiments shows that abrasion or wounding of the plant tissues is not necessary before infection can take place. That such injury would render a plant more liable to infection is, however, indicated by the

results obtained in Exps. 13 and 14.

Since the fungus grows readily as a saprophyte not only on dead violas but also on many other kinds of plants, the possibility of infection spreading to violas from various other sources must be taken into account. In this connection it may be mentioned that a culture of *M. roridum* isolated from dead vine stems was found readily to infect healthy living plants of *Viola*.

SUMMARY

Inoculation experiments with the fungus Myrothecium roridum, obtained in pure culture from dead Viola stems, are described.

A high percentage of infections was secured by applying a spore

suspension to the uninjured tissues of healthy Viola plants.

The fungus was shown to be able to infect the leaves, stems and hypocotyledonary region of growing plants.

It is concluded that *Myrothecium roridum* must be regarded as a potential parasite under natural conditions.

FOOTNOTE. After this paper had been sent to the editors a preliminary account of a serious crown rot of snapdragons caused by Myrothecium appeared in Phytopathology, xxv (1935), 969—J. J. Taubenhaus: "On a black crown rot of greenhouse snapdragons caused by Myrothecium roridum Tode."

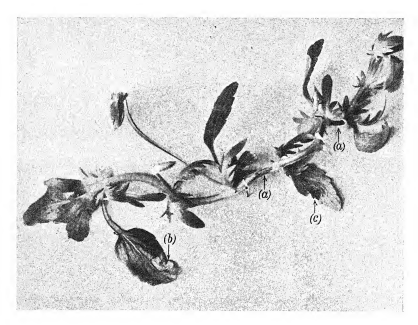


Fig. 1

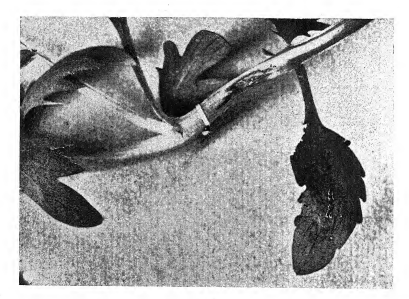


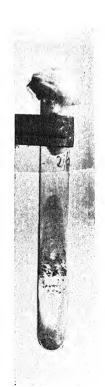
Fig. 2

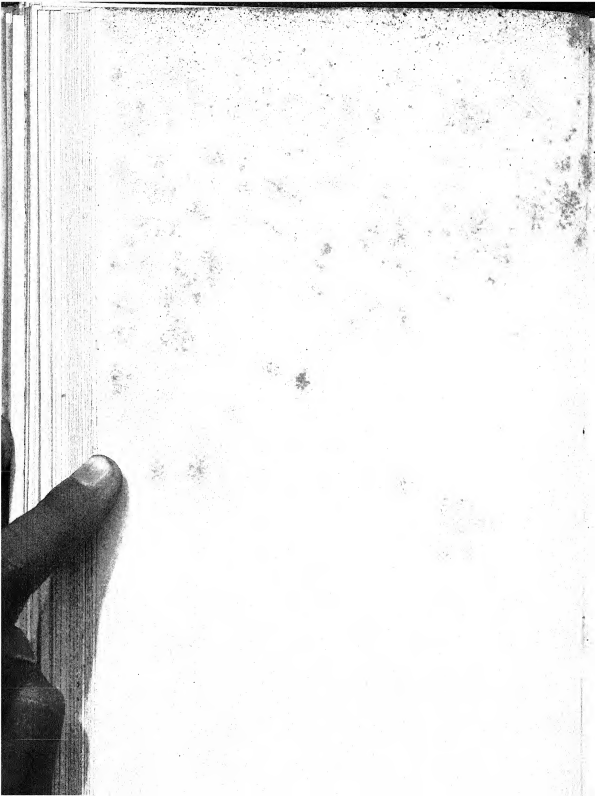




Fig. 3







EXPLANATION OF PLATES IV AND V

PLATE IV

Fig. 1. Viola seedling, "Chantryland", artificially inoculated with Myrothecium roridum showing lesions on stem (a) and on leaves (b), (c). Lesions (a) and (b) six days, lesion (c) ten days after inoculation.

Fig. 2. Part of same plant as fig. 1 enlarged showing sporodochia on the inoculated

PLATE V

Fig. 3. Viola cutting, "Maggie Mott", showing artificially produced lesions (a) on the stem, fourteen days after inoculation.

Fig. 4. Extensive lesion on stem of "Maggie Mott" Viola, fourteen days after inoculation. Note concentric wrinkling at centre of lesion and the smooth healthy internode above the one inoculated.

Fig. 5. Myrothecium roridum Tode. Three weeks' growth on cherry agar.

OBSERVATIONS

ON THE RESULTS OF INOCULATING CEREALS WITH THE SPORES OF CEREAL RUSTS WHICH DO NOT USUALLY CAUSE THEIR INFECTION

By THEODORA B. HANES, Ph.D. (CANTAB.)

(With 18 Text-figures)

INTRODUCTION

The study of specialisation among the cereal rusts was initiated by Eriksson as early as 1894, and since that time the subject has received much attention. Observations by early workers in different countries on the host range of particular rusts showed unaccountable discrepancies, which became explicable only after the recognition of numerous physiologic forms of the different rusts, the existence of which was first demonstrated in *Puccinia graminis Tritici* by Stakman and Piemeisel in 1917.

Our knowledge of the specialisation and host range of the different cereal rusts is largely based on macroscopic observations of the infected hosts, and consequently the reaction between host and parasite has been defined merely in terms of such visible symptoms as necrosis, flecking, and pustule formation. Such observations give only an incomplete picture of the development of the fungus and its relation-

ship with the host tissues.

The first careful investigations of the intimate relationship existing between host and rust were made by Marshall Ward in 1901 and 1902. Gibson found, in 1904, that chrysanthemum rust could enter various plants on which it does not normally occur. However, no true infections resulted, that is, no haustoria were formed, and by the end of the fourth day the fungus was dead. Killed host cells were observed in the vicinity of these hyphae, and Gibson suggested that the death of the hyphae was due to some poisonous substance emitted by the cells.

Marryat, in 1908, studied two varieties of wheat showing different degrees of resistance to yellow rust, *P. glumarum*. She found that the fungus entered these plants in a normal manner and produced hyphae, but development was restricted owing to the death of host tissue in the infected areas, which cut off the food supply of the parasite. In the variety "American Club", which was the less resistant, the fungus developed further but ultimately death resulted.

Stakman, in 1914 and 1915, studied the development of stem rust

of wheat on immune and susceptible varieties of that host. He was the first to study microscopically the several forms of P. graminis on hosts presenting an extreme degree of incompatibility. These studies included oats inoculated with P. graminis Tritici and P. graminis Hordei; rye, wheat, and barley inoculated with P. graminis from Dactylis glomerata; and wheat inoculated with Puccinia graminis Avenae. The experiments showed that the fungi entered "practically immune" plants in a normal manner, but after the death of a number of host cells they were unable to develop further. Stakman termed such host plants "hypersensitive". He states that immunity and resistance are independent of the state of nutrition of the host, but that the explanation probably lies in the secretion of toxins by host, or parasite, or both.

Newton, in 1922, studying *P. graminis Tritici*, showed that a resistant variety of wheat exhibited the same intolerance to the fungus as described by Stakman (20). She, however, interpreted the early cessation in the development of the fungus as being due to starvation

resulting from the death of host tissue.

More recently, Allen, in 1923 and 1926, contributed valuable information regarding the host-parasite relation by means of cytological studies of different physiologic forms of *P. graminis Tritici* on varieties of wheat showing various degrees of susceptibility and immunity. Entry of the fungus was always observed, but even in susceptible varieties the guard cells of the stomata of entry were often killed. The subsequent development of the rust varied. Khapli emmer, a variety of wheat resistant to all forms of *P. graminis*, was inoculated with forms 9, 21 and 27. The first mesophyll cell invaded always collapsed and died but the fungus survived, and occasionally produced minute uredo pustules. In addition to microscopic studies of immune and susceptible varieties of wheat inoculated with *P. graminis Tritici*, Allen, in 1926 and 1927, made cytological studies of Little Club wheat (susceptible) and Malakoff wheat (resistant) inoculated with *P. triticina* P.F. 11.

Ruttle and Fraser, in 1927, studied in detail a resistant and a susceptible variety of oats inoculated with *P. coronata* Corda. They found that the entry of the fungus was similar in both varieties. In Cowra 35, the resistant host, the first cell invaded was killed, but sometimes the fungus continued to develop and eventually produced minute pustules as recorded in *P. triticina* on Malakoff wheat (5), and *P. graminis Tritici* on Khapli emmer (3).

¹ The recent work of Forward, in 1932 (*Phytopathology*, XXII, 493–555), seems to render this view untenable, since she was able to produce changes in infection type with *P. graminis Tritici* P.F. 21 as a result of altering the host metabolism by starvation. For example, after prolonged periods of darkness, hypersensitive areas appeared on hosts which were ordinarily congenial.

The above account shows that we have little information concerning the development of cereal rusts in an extended range of plants. The experiments of Stakman (19, 20) with P. graminis apparently provide the only existing account of the development of cereal rusts on hosts on which they are not normally found in nature ("inappropriate hosts"). Most other workers have restricted their experiments to immune and susceptible varieties of the natural host.

The work now presented was designed to supplement existing knowledge concerning this aspect of the problem of specialisation

among the cereal rusts.

MATERIALS AND METHODS

Inoculation experiments were carried out with uredospores of the following rusts:

Puccinia triticina Erikss. (brown or leaf rust of wheat).

P. glumarum Tritici Erikss. (yellow or stripe rust of wheat).

P. anomala Rostrup (brown or dwarf rust of barley).

P. coronata Corda (crown rust of oats).

P. graminis Secalis Erikss. (black or stem rust of rye).

In addition, aecidiospores (P. coronata Corda) from Rhamnus catharticus and R. Frangula provided the inoculum for some experiments.

The particular physiologic forms of the fungi used in these investigations are not known, as no work had been done in England on their determination at the time these researches were carried out.

All plants used for inoculation were grown in 4-in. pots in green-houses and were from one to three weeks old at the time of inoculation. They included wheat, rye, barley, and oats, and, in some experiments, certain grasses in addition.

All the seeds showed excellent germination, except those of Agropyron repens, which proved to be unreliable. Young plants of this

grass were obtained, however, by potting pieces of rhizome.

In experimenting with hosts which were susceptible to more than one form of rust used, it was necessary to guard against accidental infection. As far as possible, therefore, experiments with different rusts were carried on in different greenhouses. Cultures of the various rusts were established on their appropriate hosts and were maintained in the greenhouses. In a few experiments spores were taken directly from the field and used for inoculation.

For easy reference the details of experiments are set out in tabular form. In these tables the cultural history of the rust used is given in an abbreviated form; for example, G 6 from wheat indicates that the particular rust has been cultured for six successive generations on wheat; G 4 from oats indicates that the rust has been cultured on

oats for four successive generations.

At the time of each experiment the germination capacity of the spores used was tested by sowing them on tap water in watch-glasses. Spores frequently showed 100 per cent. germination under these conditions, but it did not necessarily follow from this that even susceptible plants would be heavily infected.

Inoculations were made with a sterile scalpel, only the first leaf of each seedling being inoculated. All inoculated areas were marked with waterproof ink, and the plants were placed in a moist chamber

for forty-eight hours after inoculation.

In some experiments the inoculated plants were observed only macroscopically, while in others areas were removed at intervals after inoculation. These were fixed, embedded in paraffin wax (M.P. 50° C.) and sectioned. Cedarwood oil was used instead of xylol for embedding, as the material was then less brittle. Several fixatives were tried, including formol-acetic-alcohol, chrom-acetic, and chromacetic-urea of different strengths. Consistently better fixations were obtained with Allen's (2) chrom-acetic-urea, made up in the following proportions: 1 gm. acetic acid, 1 gm. chromic acid, and 0.5 gm. urea in 100 c.c. distilled water.

Sections were cut 10 \(\mu \) in thickness. For staining the following combinations were tried: Heidenhain's iron-alum haematoxylin and orange G; Flemming's triple stain; and diamant fuchsin and light green. The diamant fuchsin and light green combination was best

and was used almost exclusively.

INVESTIGATIONS

Section I. Puccinia triticina Erikss.

In 1907 Pole-Evans published an account of the histology of P. triticina. Detailed cytological studies of this rust both on a susceptible and a resistant variety of wheat were made by Allen in

1926 and 1927.

P. triticina was collected at the University Farm, Cambridge, on October 11, 1927. The first transfer was made to seedlings of Wilhelmina wheat in the greenhouse and an excellent crop of spores developed. By successive transfers to Wilhelmina wheat the culture

was maintained for two and a half years.

The experiments with P. triticina were more numerous than those with other rust forms, since an abundance of spores from greenhouse cultures was always available. Inoculations were carried out on wheat, rye, barley, and oats. The varieties used were Wilhelmina, and occasionally Persian Black wheat; rye of unknown variety; Spratt Archer barley; and Grey Winter oats.

(1) Experiments for macroscopic study of inoculated plants

Inoculations were made on wheat, rye, barley, and oats, but no inoculated areas were fixed. Daily observations were made on all plants. Thirty-five experiments were made, of which twelve repre-

sentative ones are described in Table I.

Wheat. In the thirty-six experiments in which wheat seedlings were inoculated with spores of P. triticina heavy crops of spores resulted in thirty-three; in all but one of these, all the seedlings inoculated were infected. In three experiments no infection resulted, and in two of these the failure of infection was correlated with very low germination of the inoculum.

Rye. In the thirty-four experiments in which rye seedlings were inoculated, macroscopic observation showed that infections resulted in all but five. There was, however, a striking variation in the type of infection from one experiment to another. These experiments fall

roughly into the following classes:

(i) Normal infection, as in wheat, resulting in a heavy crop of spores, and without necrosis. Five experiments, in which fifty-six out of fifty-eight inoculated plants were heavily infected.

(2) Weak infection resulting in small pustules with no necrosis. Three experiments in which only four out of twenty-seven plants

produced pustules.

(3) Mixed infections in which some leaves produced heavy crops of spores with no necrosis, while others produced few or no spores and showed considerable necrosis. Four experiments in which twenty-nine out of forty plants produced pustules.

(4) Infections resulting in minute pustules on a few leaves accompanied by necrosis, and large necrotic areas on most of the inoculated

leaves.

(5) Infections resulting in no pustules but considerable necrosis. Consideration of the data concerning these experiments has failed to reveal any clue as to the nature of the factors which govern the type of infection on rye.

Barley. In the twelve experiments in which barley seedlings were inoculated, weak infections resulting in minute pustules occurred in

three. There was no necrosis on barley.

The total number of barley plants inoculated in these experiments

was III, and of these ten became weakly infected.

In the single experiment (No. 10) in which spores produced on barley were used as the inoculum, no infection resulted on barley seedlings.

Oats. In the eleven experiments in which 111 oat seedlings were inoculated no definite sign of infection was observed, although one

plant showed flecking.

Table I. Puccinia triticina. Experiments for macroscopic study

				Teresiste Jor macroscopio stataj
xp. Vo.	Date	Plants inoculated	Culture used	Results
I	Jan. 9, 1928	Wheat* Rye Barley Oats	G 2 from wheat Germ. 100 %	Wheat 10/10,† good infection Rye 1/10, weak infection Barley 0/10 Oats 0/10
2	Feb. 25, 1928	Wheat Rye	G 3 from wheat Germ. 100 %	Wheat 12/12, good infection Rye 1/12, weak infection. Large necrotic patches on many leaves of rye
3	June 15, 1928	Wheat Rye	G 7 from wheat Germ. 50 %	Wheat 20/20, good infection Rye 0/20 (no necrosis)
4	Aug. 6, 1928	Wheat Rye	G I from Wil. taken from P.B. wheat Germ. 78 %	Wheat 16/16, good infection Rye 20/20, good infection. Some rye leaves quite as heavily infected as wheat. No sign of necrosis on rye but a few plants were more definitely flecked than wheat. The flecks were normal in appearance
5	Aug. 28, 1928	Wheat (P.B.) (Wil.) Rye Barley Oats	G 4 from P.B. Germ 75 %	Wheat (P.B.) 10/10, good infection; (Wil.) 10/10, good infection Rye 2/10, poor. Large necrotic patches on some inoculated leaves. Barley 1/10, weak infection. A few minute scattered pustules; first positive infection of barley with P. triticina Oats 0/10
6	June 11, 1929	Wheat Rye	G 12 from wheat Germ. 100 %	Wheat 12/12, good infection Rye 12/12, good infection. Slight necrosis. Pus- tules on both hosts 7 days after inoculation
7.	June 26, 1929	Wheat Rye Barley Oats	G 13 from wheat Germ. 100 %	All plants showed definite flecking on 6th day Wheat 20/20, good infection Rye 20/20, good infection, no necrosis on rye Barley, oats, no pustules
8	July 14, 1929	Wheat Rye	G 14 from wheat Germ. 100 %	Wheat 10/10, good infection Rye 0/10, necrosis
9	July 19, 1929	Wheat Rye Oats Barley	G 14 from wheat Germ. 20 %	Wheat 10/10, good infection Rye 5/10, weak infection, necrosis Barley 5/10, weak infection, very small pustules but no necrosis Oats 0/10
10	Aug. 2, 1929	Barley	G I from barley (transferred from wheat), very few spores available	No sign of infection after 3 weeks
11	Aug. 7, 1929	Wheat Rye	G 15 from wheat Germ. 100 %	Wheat 10/10, good infection Rye 10/10, good infection. 5 leaves produced large pustules and infection appeared normal: on the other 5, pustules were smaller and there was some necrosis
12	Aug. 25, 1929	Wheat Rye Barley Oats	G 16 from wheat Germ. 100 %	Wheat 9/9, good infection Rye 6/6, good infection; smaller pustules but no necrosis Barley 4/10, very minute pustules but no necrosis Oats 0/10

^{*} Wheat: unless Persian Black (P.B.) is specified, the variety used was Wilhelmina (Wil.).
† The denominator of the fraction represents the number of plants inoculated, the numerator the number which produced pustules. Unless stated there was no sign of flecking on the plants which did not produce pustules.

(2) Experiments for microscopic study of inoculated plants

These experiments were carried out in order to follow microscopically the development of *P. triticina* in wheat, rye, barley, and oats. Seedlings were inoculated as before, and marked areas were removed at intervals and prepared for microscopic examination. The varieties used were the same as before, only one variety of wheat (Wilhelmina) being used for microscopic study.

The details of the experiments are given in Table II.

Table II. Puccinia triticina. Experiments for microscopic study

	Tab	de II. Pu	iccinia trii	icina.	Experiments for interescopie study
				Times of fixations: days after	
Exp. No.	Date	Plants inoculated	Culture used	inocula- tion	
36	Jan. 9, 1928	Wheat Rye Barley Oats	G 2 from Wil. wheat	5 7∫	Flecking on wheat only
				10 21	Pustules on wheat only Pustules on wheat only. Necrotic patches on rye which had appeared as small flecks on the 13th day. No sign of infection on barley or oats
37	Feb. 25, 1928	Wheat Rye	G 3 from Wil. wheat	17	Numerous pustules on wheat. Definite necrosis on inoculated leaves of rye, and a few minute pustules on one leaf
38	June 17, 1929	Wheat Rye	G 11 from Wil. wheat	3	Flecking on wheat and rye
				4 8	Numerous pustules on wheat. Numerous pustules on some leaves of rye, but fewer and smaller pustules on others accompanied by necrotic patches
39	June 26, 1929	Wheat Rye Barley Oats	G 13 from Wil. wheat	10	Numerous pustules on wheat and rye. No pustules on barley and oats, but definite flecking
40	July 1, 1929	Wheat Barley Oats	G 13 from Wil. wheat	3) 4 6	No flecking Flecks on wheat only Numerous pustules on wheat. No sign of infection
				10	on barley or oats
41	Sept. 28, 1929	Wheat Rye Barley Oats	G 19 from Wil. whea	1 2 4 6	No flecking Faint flecking on wheat only Definite flecking on wheat only Wheat as above. A few flecks on rye, but none on barley or oats
				8 11	No sign of pustules. A few faint flecks on barley Numerous pustules on wheat. Necrotic patches on rye; no pustules Numerous pustules on wheat. A few small pustules

on rye and large necrotic patches. A few minute pustules on barley and no sign of necrosis. Very faint flecking on oats but no sign of pustules The data given in the last column of the Table show that the rate of development of *P. triticina* in these plants varied from one experiment to another. The following accounts of the development of the fungus are generalised from the different experiments, but variations in the developmental rate will be indicated.

A study was first made of the progress of the fungus in wheat, the original host. This study provided a basis for comparing its progress

in rye, barley and oats.

Development of Puccinia triticina in wheat

During the first twenty-four hours many of the spores germinated. Appressoria were formed over the stomata, and the contents of these passed into vesicles in the substomatal cavities. The stomata of entry were not damaged by the fungus. By the second day infecting hyphae were seen growing toward the mesophyll cells, and on the third day

many had formed haustoria.

Fig. 1 (three-day material) shows an entry which is typical of those found in wheat. Here the vesicle is almost empty. The infecting hypha branched into two shortly after its formation. Both branches were closely applied to the walls of mesophyll cells. The first haustorial mother cell (now empty) was cut off at the tip of branch a. This produced haustorium d, which lay in close contact with the host cell nucleus e. Hypha f arose just behind the haustorial mother cell. It was full of contents and lay across the invaded host cell. Branch b of

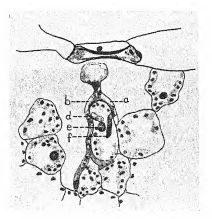


Fig. 1. (×315,)

the infecting hypha was much longer than a, but had formed no haustoria b at the time of fixation. Hypha b was full of contents and showed very clearly the groups of three nuclei which are characteristic of the mycelium of P. triticina as noted by Allen (4). There was no disturbance of the surrounding tissues.

After establishing contact with the host tissue by the formation of the first haustorium, the fungus grew rapidly. New hyphae and haustoria continued to develop, and by the fourth day an intimate relation was established with the host. From the seventh day dense wefts of hyphae, in preparation for pustule formation, occurred at intervals below the upper and lower epidermis. Even at this stage the host cells retained their normal appearance. By the eighth day

many pustules had developed. Hyphae in the tissue below the pustules were empty and frequently septate, and large haustoria were present in the host cells. Mesophyll cells in the pustule region were still living, and retained their contents. As noted by Allen (4), a few dead host cells were occasionally found in the vicinity of pustules, apparently crushed by the great mass of hyphae formed.

The development of *P. triticina* on Wilhelmina wheat has been described briefly, since its development on this host agrees closely with that described by Allen for the rust on Little Club wheat, a

susceptible variety.

Development of Puccinia triticina in rye

Different types of reaction may occur between rye and *P. triticina*. This is clear from the macroscopic observations recorded in Table II. Different types of reaction might occur on different leaves of the

same plant, or on different parts of the same leaf.

The variability in the type of infection in rye became evident from the macroscopic observations, some of which are set out in Table I; microscopic study serves to distinguish four different types of reaction in rye, A, B, C and D. These reaction types showed differences in the development attained by the fungus, its relation with the host tissue, and the final fate of the intercellular mycelium.

Type A reaction in rye.

The spores germinated, formed appressoria, and many of the fungi entered the leaves during the first twenty-four hours without damaging the stomata. Vesicles with their infecting hyphae were numerous in the substomatal cavities, and by the second day some of these had formed haustoria in host cells.

Fig. 2 (from two-day material) shows entry into rye at this time.

Two empty appressoria can be seen outside a stoma, and two vesicles with their infecting hyphae in the substomatal cavity. Infecting hypha a has already formed a haustorium d, which lies in contact with a host nucleus, and the terminal haustorial mother cell is now empty. Other infecting hyphae were sometimes branched. No

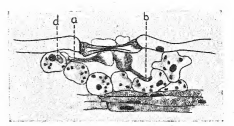


Fig. 2. (×315.)

harm was done to the mesophyll cells.

In one leaf area of rye fixed on the second day, the fungus had already developed several healthy-looking hyphae, a stage of development not usually reached in wheat until the third or fourth day. In some leaf areas (Exp. No. 38) the fungus was well established by the fourth day, and its relation with the host appeared to be quite

congenial.

By the seventh day the intercellular hyphae had spread considerably and formed wefts of mycelium beneath the upper and the lower epidermis. The hyphae were now mostly empty and septate, and large haustoria were seen in the host cells. By the eighth day many pustules had formed. In Exp. No. 38 these were, in general, smaller than those on wheat, but otherwise similar. In Exp. No. 39 large pustules were produced on both surfaces of the leaves.

In type A reaction, the development of *P. triticina* in rye is the same as in wheat, the relation between host and parasite remaining

congenial throughout.

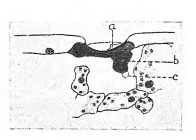


Fig. 3. $(\times 315.)$

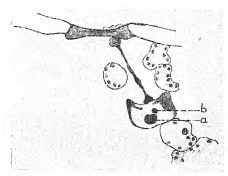


Fig. 4. (×315.)

Type B reaction in rye.

An examination of leaf areas from several experiments showed that sometimes *P. triticina* made only restricted progress in rye, in contrast to the development which has been described under type A reaction.

The fungus entered as before, but the guard cells of the stomata of entry were usually dead and discoloured. Many vesicles in material fixed after twenty-four hours retained their contents, but these were frequently deeply stained and no infecting hyphae had developed.

Fig. 3 (twenty-four-hour material) shows one of these vesicles. The guard cell, seen in section, is dead, and the shrivelled remains of the appressorium a are outside. The vesicle b is dead, and very deeply stained. The projection at c is probably the rudiment of an infecting hypha.

For three weeks after inoculation, vesicles were seen in the substomatal cavities. Some produced infecting hyphae, and occasionally

a haustorium was developed.

A later entry is shown in Fig. 4 (from seventeen-day material).

Here the infecting hypha produced a haustorium a in the first host cell encountered. This cell (nucleus, b) is now dead and the fungus is

considerably shrivelled.

These figures illustrate the maximum development of the fungus in type B reaction. The fungus did not succeed in establishing itself in the host tissue; there was no development after the formation of the infecting hypha, which only occasionally produced a haustorium. It was clear that in this type of reaction an antagonistic relation existed between the fungus and host. Only slight damage was done to the host tissue, since at most only two or three host cells were killed.

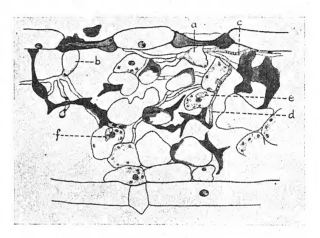


Fig. 5. (×315.)

Type C reaction in rye.

In this type of reaction, as in type B, the relation between rust and host was definitely antagonistic. The fungus, although it did not develop so far as to produce pustules, made considerably more progress than in type B reaction, and consequently larger areas of leaf

tissue were damaged, forming visible necrotic patches.

In leaf areas where type C reaction was observed many of the entries showed the type B reaction. Some entries, however, succeeded in establishing an intercellular mycelium, and the vesicles with their infecting hyphae were found empty in the substomatal cavities. If now the mycelium developed, type C reaction was encountered. The characteristics of this type of reaction are shown in Fig. 5 (eleven-day material), where two separate entries are seen at neighbouring stomata. The guard cells are dead. The vesicles a and b are practically empty. A few intercellular hyphae resulting from these entries can be seen, some (a and b retaining their contents while others are

empty. Haustoria can be seen at e and f, but very few host cells are living. Some are shrunken and so deeply stained that it is impossible to distinguish the contents, while others are empty. Similar empty cells were found by Allen (5), when Malakoff, a resistant variety of wheat, became infected with P. triticina P.F. 11. Such infected zones of leaf tissue are local and are separated by zones of healthy cells. The abundance of non-functioning haustorial mother cells is a striking feature of many of the zones exhibiting the type C reaction.

In type C reaction antagonism existed between the fungus and rye, damage done to the host involving considerable areas of leaf tissue. The condition of the fungus itself was weak, and there were no signs of pustules.

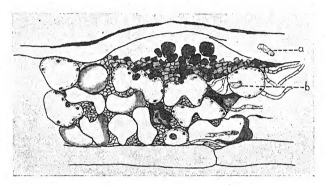


Fig. 6. $(\times 315.)$

Type D reaction in rye.

Type D reaction resembles most closely type C, but differs from it in that the fungus succeeded in producing a few minute pustules in the necrotic areas. These appeared in Exp. No. 41, for example, sixteen days after inoculation, whereas pustules appeared on wheat on the eleventh day.

The relation between rye and the rust in type D reaction is illustrated in Fig. 6 (seventeen-day material). Some host cells are empty; others are shrunken and have deeply stained contents; still others are characterised by irregular thickenings of their walls. (Such thickenings were also observed in type C reaction.) They correspond closely to the "swellings" and "warts" described by Allen (5) as occurring outside the infected zones of Malakoff wheat. In rye, however, these swellings were found directly in the infected areas. Fragments of hyphae seen in the tissue have sparse, vacuolated contents, or none at all. Very few haustoria were formed; two are shown, one in an epidermal cell at a, and one in a mesophyll cell at b. Although small pustules were formed, few spores reached maturity.

Development of P. triticina in barley

The rust enters barley as freely as it does wheat and rye. Actual counts were not made, but by the end of the first day vesicles were numerous in the substomatal cavities and the stomata were not damaged. New vesicles continued to develop during the first eight days after inoculation. The fate of most of these became clear from a study of later fixations in which numerous dead and shrivelled vesicles with infecting hyphae were often found in the substomatal cavities, death having occurred before relation was established with the host. These entries were similar to some in type B reaction in rye, except that in barley the guard cells were not killed (cf. Fig. 4).

Barley fixed seventeen days after inoculation in Exp. No. 41 showed zonal infection similar to that described in type D reaction in rye. In barley there were fewer dead cells, indicating that the reaction in barley was less violent than in rye. Only rarely did a few minute pustules develop in barley, and these were considerably aborted. On the rare occasions where *P. triticina* reached the reproductive stage in barley, there was an antagonistic relation between barley

and the rust.

Development of Puccinia triticina in oats

The rust entered oats freely, forming numerous vesicles in the substomatal chambers. The stomata were not damaged. Frequently two or more vesicles were found in the same cavity, and occasionally

infecting hyphae were found.

Some entries seen in oats showed disturbance of the surrounding mesophyll cells although no haustoria had been formed. The rust made little progress in oats, and only once was a small haustorium seen, marking the most advanced stage that the fungus achieved in oats. Leaf areas fixed eight, eleven, and seventeen days after inoculation showed no further development.

DISCUSSION OF EXPERIMENTS WITH PUCCINIA TRITICINA

It seems clear from these experiments that *P. triticina* (in England) is not sharply specialised to wheat. Eriksson (8) found that rye could sometimes be infected with this rust, and Mehta (15), working in England, obtained successful infection on three out of twenty-one seedlings of rye inoculated. These authors were in agreement that rye was subject to "casual infection" by *P. triticina*.

Different types of reaction result when rye is inoculated with *P. triticina*. The phenomena encountered suggest a similarity to the "heterogeneous or X-reaction" described by Stakman and Levine (21) for the types of reaction observed on some wheat varieties inoculated

with certain biologic forms of P. graminis Tritici.

A further indication that *P. triticina* is not closely specialised to wheat was obtained from Exps. Nos. 5, 9, 12 and 41 (Tables I and II). In these minute pustules were produced on barley.¹ Microscopic examination of inoculated areas removed during the course of Exp. No. 41 revealed infected zones similar to those of type D reaction in rye. Dead mesophyll cells were less numerous in barley, indicating that the reaction was less violent than in rye. The pustules which developed were small and aborted.

Oats was consistently immune to *P. triticina*. Flecking was observed in three out of fifteen inoculations, but no pustules developed. Microscopic examination showed that the rust frequently entered oats, but thereafter made little progress; only one small haustorium was found

in the many entries which were examined.

One further point deserves comment. Experiments were carried out to determine whether a sojourn of the rust on rye and barley would increase its virulence on these respective hosts. Spores produced by successful infections on rye with *P. triticina* were used to inoculate seedlings of rye and wheat. The results showed that the virulence was not increased, since these inoculations produced only weak infections on rye accompanied by necrosis, whereas heavy crops of spores were produced on wheat in every trial. Rye was apparently more susceptible to *P. triticina* taken directly from wheat. On the one occasion when pustules on barley produced sufficient spores to be used as an inoculum, these were used to inoculate barley seedlings (Exp. No. 10). No infections resulted."

Section II. Puccinia glumarum Tritici Erikss.

P. glumarum Tritici is one of five specialised forms of P. glumarum described by Eriksson in 1894. Forms of yellow rust occur also on barley, rye, and various grasses. According to Eriksson, the rust on wheat is a specialised form for that host, and does not infect barley or rye. In the United States, however, Hungerford and Owens (12) found that P. glumarum Tritici would infect rye "moderately", barley "slightly", and forty-seven wild grasses. They found that oats did not become infected when inoculated with this form.

Mehta (15) showed the strict specialisation of P. glumarum Tritici to

wheat; barley and rye were not infected by this form.

The histology of *P. glumarum Tritici* has been studied by Pole-Evans (17) and by Marryat (14), and in 1928 Allen published an account of the cytology of *P. glumarum* on *Bromus marginatus* and *Triticum vulgare*.

¹ Prof. H. S. Jackson has informed me that *P. triticina* has been found in the United States, occurring naturally on a wild variety of barley.

ESTABLISHMENT OF THE CULTURES

Norka wheat, a variety very susceptible to yellow rust, was used exclusively for the greenhouse cultivation of this rust. Plants of Norka wheat kept on the roof of the Botany School, Cambridge, became infected with yellow rust in October 1927. Successive transfers of the rust were made to Norka wheat in the greenhouse, and for eight months the culture was maintained. In the hot summer of 1928, however, the culture was lost at the end of June. Infected leaves of Little Joss wheat, found at the University Farm on June 30, 1929, provided spores from which another culture was established and maintained in the greenhouse for six months.

EXPERIMENTS WITH PUCCINIA GLUMARUM TRITICI

Seedlings of Norka wheat, Grey Winter oats, rye (variety unknown), and Spratt Archer barley were inoculated with spores of P. glumarum Tritici.

The experiments and results are set out in Table III.

Table III. Experiments with Puccinia glumarum Tritici

Exp. No.	Date	Plants inoculated	Culture used	Results
I	May 21, 1928	Wheat Rye Barley Oats	G 4 from Norka	Flecking on wheat on 7th day. Pus- tules on wheat only on the 10th day. Brownish yellow streaks on barley 14th day, but no indication of in- fection on oats or rye
2	July 15, 1929	Wheat Rye Barley Oats	G I from Norka (new culture)	Flecking on wheat on 7th day. Very small pustules on wheat only on 12th day. No sign of infection on barley, oats or rye
3	Oct. 6, 1929	As in 1 and 2	G 3 from Norka	Moderately good infection on wheat after an incubation period of 3 weeks. No sign of infection on barley, oats, or rye
4	Oct. 30, 1929	As in 1 and 2	G 4 from Norka	As in Exp. No. 3 above
5	Nov. 26, 1929	As in 1 and 2	G 3 from Norka	As in Exp. No. 3 above

In Exps. Nos. 1 and 2, inoculated areas of wheat, rye, barley and oats were removed at intervals after inoculation and prepared for microscopic study. The remaining experiments, Nos. 3, 4 and 5, were carried out for macroscopic observation only.

It seemed clear from the macroscopic observations in the experiments of Table III, that yellow rust of wheat is closely specialised to that host. This was confirmed by the microscopic study of the inocu-

lated plants.

Development of Puccinia glumarum in wheat

Within twenty-four hours many spores germinated and formed long germ tubes on the leaves. No entries were found until the second day, although Allen (6) found entries sixteen hours after inoculation; she noted, however, that entry might be delayed for several days. This was also found in my experiments, where new vesicles were found in the substomatal cavities as late as the seventh day. The fungus entered without forming an appressorium, as pointed out also by Pole-Evans (17), and the stomata of entry were not damaged. The germ tubes usually entered at one end of a stoma, but were occasionally found near the centre.

The prolonged initial developmental period of *P. glumarum* under certain conditions is well known. By the fourth day many infections showed no advance beyond the formation of the first haustorium from the infecting hypha; other infections, however, made considerable progress, and thick, non-septate hyphae with numerous

nuclei were found among the mesophyll cells.

Leaf areas fixed on the seventh day showed a marked increase in mycelial development. (This coincides with the appearance of flecking on this host.) Hyphae were both branched and unbranched, and they were full of contents. The long, runner-like hyphae described by Allen (6) were frequently seen in leaf areas fixed on the seventh day.

By the tenth day the hyphae were narrower and their contents less dense. Haustoria at this time were mostly of the "hammerhead" type and were found in both epidermal and mesophyll cells. Small pustules had sometimes developed by this time; the epidermis was still unruptured over them, and the hyphae in the pustule regions

were considerably drained or else quite empty.

On the fourteenth day numerous pustules had formed on both surfaces of the leaves, and the underlying mesophyll tissue was often almost completely obliterated by the profuse development of hyphae. These were empty and septate. Where the hyphae were less dense, haustoria, large and of various shapes, were present in the host cells. These completely replaced the "hammerhead" type, which were so common on the tenth day. The host cells were impoverished but still living.

Development of Puccinia glumarum Tritici in rye

No entries were found in rye until the fourteenth day after inoculation. Earlier material showed many spores of the inoculum on the surface of the leaves; some of these had germinated and formed long germ tubes, which had shrivelled, however, without entering.

In material fixed on the fourteenth day several entries were seen.

The fungus, however, had made little progress and was invariably dead. The guard cells were also dead and discoloured. Fig. 7 (fourteen-day material) shows one of these entries. The empty

spore, somewhat shrivelled, lies near the stoma where the fungus entered. As usual with P. glumarum, entry was made at one end of the stoma. The guard cellseen in section is dead and discoloured, and closely applied to it is the vesicle a, typical of this rust, now empty. In this entry two large infecting hyphae, b and c, developed. Hypha b, soon after its formation, attacked and killed a mesophyll cell bordering on the substomatal cavity. The hypha is now misshapen and deeply stained. The empty haustorial mother cell d is seen outside the dead host cell, and it appears that the large haus-

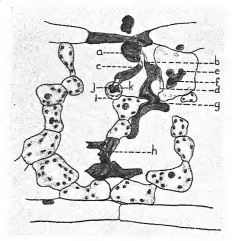


Fig. 7. (×315.)

torium e coiled itself about the host nucleus f. Both haustorium and nucleus are dead and deeply stained, and in the cell the remains of a few chloroplasts are faintly visible. The hypha is considerably swollen behind the empty haustorial mother cell, a fact which suggests that some food from the host had passed back through the haustorial mother cell before death occurred. After this failure to establish contact with the host, hypha b put out a branch g, but no infection resulted. This branch g is dead and deeply stained like the rest of the fungus. The two host cells in contact with g are living and retain their normal appearance. An adjacent cell h, though not in contact with the fungus, is dead and deeply stained. The other infecting hypha c branched into two just before meeting the host cell i, which borders on the substomatal chamber. Both branches of c produced haustoria in this cell. It appears that these haustoria, j and k, arose without the usual formation of haustorial mother cells. Both haustoria are dead, and lie in contact with the nucleus of the dead host cell.

This entry represents the greatest progress made by *P. glumarum Tritici* in rye; the fungus was short-lived, and only a few host cells were killed.

Microscopic study showed that rye is immune to yellow rust of wheat. For some time after inoculation, although the spores germinated on the leaves, they failed to effect entries. After this delay, however, entries were made, but these were not nearly so numerous as in wheat. Only slight progress was made after entry, the infecting hyphae producing at most one or two haustoria before death. On the rare occasions when this development was attained, a few host cells were killed. The relation between rye and *P. glumarum Tritici* was uncongenial.

Development of Puccinia glumarum Tritici in barley

The fungus was first found in leaf areas of barley fixed on the seventh day. As in rye, the guard cells of the stomata of entry were always dead and discoloured. Entries were common in the seven-day material of barley; infecting hyphae had developed but no haustoria had been formed, and in all cases the fungus was dead.

A further stage of development in barley is seen in Fig. 8 (fourteen-day material). The guard cell is dead and the vesicle quite empty.

As before, the fungus is now dead, but several host cells were affected before death occurred. Two infecting hyphae developed from the vesicle. a and b (hypha b has been broken in sectioning). Hypha a formed a large haustorium d in the mesophyll cell c, which is devoid of contents. The haustorium has a thick neck; its mother cell e, now empty, is closely applied to the wall of the invaded host cell. An adjacent mesophyll cell shows a distinct thickening of its wall. The other infecting hypha b extended further but did not produce a haustorium. It penetrated some distance into the leaftissues, coming into contact with several mesophyll cells. These developed thickened walls where they touched the hypha, but they were still

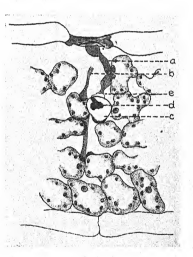


Fig. 8. (×315.)

living. Damage done by the fungus was accordingly slight, the area of tissue affected being small.

Other leaf areas of barley taken fourteen days after inoculation showed yellow streaks at this time. It was found that the rust had made considerably more progress in these areas. Thick intercellular hyphae were often found amongst the mesophyll cells, but haustoria were rare.

A section through one of these areas is drawn in Fig. 9 (fourteen-

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day material). Hypha a, which still retains most of its contents, formed a haustorium at c, which lies in contact with the cell nucleus. The empty haustorial mother cell is closely applied to the wall of the invaded cell. Hypha d is dead and its contents deeply stained. Some host cells in contact with these hyphae are devoid of contents.

No extensive damage was done by the hyphae in these streaked leaf areas. At most, groups of five or six dead host cells were found. This was the most advanced stage reached by the fungus in barley.

Development of Puccinia glumarum Tritici in oats

Entries in oats were less numerous than in rye and barley. As in rye, no entry was seen until several days after inoculation. The guard cells of the stomata of entry were always killed, and the infecting hyphae quickly died.

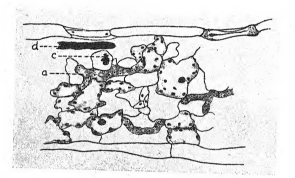


Fig. 9. (×315.)

DISCUSSION OF EXPERIMENTS WITH PUCCINIA GLUMARUM TRITICI

The development of yellow rust in Norka wheat, except for slight points, is in close agreement with the published accounts of Pole-Evans (17), Marrayat (14), and Allen (6). The "intracellular bodies" described by Allen were not found in the infected leaves of Norka. A congenial relation existed between this wheat and the rust; only in regions of heavy spore production were the host cells harmed.

Rye, barley and oats were immune from *P. glumarum Tritici*. Only in one experiment (No. 1) was there any outward sign of infection, some of the inoculated areas of barley becoming streaked.

In these streaked areas in barley the rust had produced thick intercellular hyphae in the mesophyll, but little damage was done to the leaf although some cells were killed and devoid of contents. The reaction between barley and the rust was not violent, and quite

unlike that in the immune wheats such as Einkorn or American Club, studied by Marryat. Usually, the rust made little progress in barley after entry. No relation was established with the mesophyll cells in some infections, while in others slight growth took place after the formation of the first haustorium. The walls of mesophyll cells were always considerably thickened where they came in contact with hyphae.

The rust succeeded also in entering rye and oats; as in barley, the guard cells of the stomata of entry were killed. In rye and oats, however, the mycelia were always short-lived. In rye there was no progress beyond the occasional formation of haustoria, while in oats no haustoria were seen. Damage done to the tissues of these plants was negligible, since only a small number of dead mesophyll cells

were found near the points of entry.

Section III.

Puccinia anomala Rostrup=Puccinia simplex Erikss. & Henn.

Mehta (15) inoculated wheat, rye, barley and Agropyron repens with brown rust of barley. He found that the rust attacked only barley. In studying the host specialisation of Puccinia anomala Mains (13), in extensive inoculations of grasses, succeeded in infecting only five closely related species of Hordeum. Pole-Evans (17) described briefly the histology of this rust on barley.

ESTABLISHMENT OF THE CULTURE

The culture was established on Spratt Archer barley in the greenhouse from spores collected on the University Farm, Cambridge, on May 11, 1928. The culture was maintained on this variety for nearly two years.

EXPERIMENTS WITH PUCCINIA ANOMALA

Spores of this rust were used to inoculate Spratt Archer barley, Grey Winter oats, Wilhelmina wheat, and rye of unknown variety.

The experiments are set out in Table IV.

In Exps. Nos. 1 and 4 inoculated leaf areas of all plants were fixed at one, two, three, six and nine days, respectively, after inoculation. In addition, areas of wheat and barley from Exps. Nos. 2 and 5 were fixed fourteen days after inoculation, although these experiments were originally intended for macroscopic observation only. In the other experiments the inoculated plants were kept under observation for one month, no leaf areas being removed.

Table IV. Experiments with Puccinia anomala

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Exp. No.	Date	Plants inoculated	Culture used	Results
I	July 10, 1928	Barley Wheat Oats Rye	G 3 from barley	Pustules only on barley, after 7 days. No sign of infection on the other plants
2	Sept. 5, 1929	As in 1	G 2 from barley	Faint flecking on barley and wheat after 3 days. Numerous pustules on barley only on 6th day. Wheat definitely flecked after 21 days, but no sign of infection on rye and oats
3	Sept. 9,	Barley Wheat	G 2 from barley	Numerous pustules on barley after 8 days. No sign of infection on wheat at end of 30 days
4	Sept. 17, 1929	Barley Wheat Oats Rye	G 3 from barley	Numerous pustules on barley after 8 days. No infection of the other plants at the end of 30 days
5	Sept. 30, 1929	As in 4	G 4 from barley	Numerous pustules on barley after 10 days. Faint flecks on wheat on 7th day; but no further infection after 30 days. No infection on oats or rye
6	Oct. 15, 1929	As in 4	G 5 from barley	Flecks on barley and wheat on 3rd day. Numerous pustules on barley only, on 8th day. No further infection on wheat, and no sign of infection on rye and oats after 30 days
7	Oct. 28, 1929	As in 4	G 6 from barley	Flecks on barley and wheat on 8th day. Pustules on barley only, on 11th day. No further infection on wheat, and no signs of infection on rye and oats after 30 days
8	Nov. 16, 1929	As in 4	G 7 from barley	On 11th day—numerous pustules on barley with- out previous flecking, but wheat flecked at this time. No further infection on wheat, and no signs of infection on oats or rye after 30 days
9	Nov. 23, 1929	Barley Wheat	G 7 from barley	Fair infection on barley, pustules appearing on 17th day. No sign of infection on wheat after 30 days

Development of Puccinia anomala in barley

By the end of the first day most of the spores had germinated and several entries were found. Germ tubes were occasionally seen which had branched into two on the leaf surfaces. Appressoria were formed over the stomata, and the contents of these passed into narrow elongated vesicles in the substomatal cavities, lying parallel to the guard cells. Infecting hyphae were not seen until the second day; these developed from one or both ends of the long vesicles.

By the end of the third day the fungus had made considerable progress. Delicate intercellular hyphae ramified in the host tissue, and numerous haustoria were present in the mesophyll cells. The infected zones were more or less local, as they were in material fixed six days after inoculation. Occasionally long, runner-like hyphae passed from one infected zone to another, often encountering as many as twelve cells before forming a haustorium. Some pieces of barley

leaf fixed on the sixth day showed various stages in reproductive activity. Wefts of mycelium in preparation for pustule formation were found under the epidermis, and pustules were sometimes present on both surfaces of the leaves. Hyphae in the pustule regions were empty and septate, and numerous uninucleate haustoria, more or less drained, lay in the host cells. The host cells retained their living contents. Throughout the development of the rust the relation was congenial.

Development of Puccinia anomala in wheat

The leaf areas of wheat fixed at intervals in Exps. Nos. 1 and 4 showed no entry of the fungus, in spite of the fact that many of these areas were flecked at the time of fixation. It seems clear that flecking here was not due to the fungus. Many spores had germinated on the leaves, but the germ tubes shrivelled without forming appressoria.

In Exps. Nos. 2 and 5, however, in which flecking of wheat was also observed, areas removed fourteen days after inoculation showed entry of the fungus. Where entry had occurred the guard cells were always dead. Sometimes no infecting hyphae were developed from the vesicles, but frequently one or two were found. The vesicles and infecting hyphae were always quite empty and much shrivelled. Only once was a small haustorium seen.

However, rather large groups of cells were sometimes killed around points of entry. Mycelium could not be seen in these dead areas, nor could haustoria be distinguished, as the cells were shrunken and deeply stained. The empty vesicles found in the substomatal chambers alone suggested that the damage to the leaf tissue was caused

by the fungus.

Development of Puccinia anomala in oats

Germinating spores were seen on the surface of the leaves at all times of fixation, that is up to the ninth day after inoculation, but only occasionally was entry achieved. When this occurred the guard cells were killed. No infecting hyphae developed and the mesophyll cells around the vesicles appeared normal.

Development of Puccinia anomala in rye

A few entries were observed on the second day after inoculation. These were similar to entries found in oats at this time. The guard cells of the stomata of entry were always killed. No infecting hyphae developed from the vesicles, and the surrounding mesophyll tissue showed no signs of disturbance. No further progress was made in rye.

DISCUSSION OF EXPERIMENTS WITH PUCCINIA ANOMALA

The experiments indicate a close specialisation of the rust to barley, a congenial relation being maintained between barley and the rust throughout its development.

P. anomala occasionally entered wheat, oats and rye, and when this occurred the guard cells of the stomata of entry were always killed. Only wheat, among these plants, showed flecking, but this also occurred on some leaves in which entries had not been made. A single small haustorium was found in wheat, and here the invaded host cell was not killed. Mesophyll cells in contact with infecting hyphae had thickened walls. In some entries in wheat it was clear that a violent reaction had taken place resulting in the death of groups of mesophyll cells, in addition to the guard cells. Hyphae could not be seen amongst these dead tissues surrounding such entries. The damage done to wheat was never extensive.

Entries were fewer in oats and rye than in wheat. No infecting hyphae developed, and no damage, apart from the death of guard cells, resulted. Not a single dead mesophyll cell was seen in oats or rye.

Section IV. Puccinia coronata Corda

In 1922 Hoerner carried out extensive inoculation experiments on barley, rye, wheat, and many grasses, using crown rust of oats from different localities. The results showed that crown rust of oats in the United States has an extensive host range under greenhouse conditions, since, in addition to successful infections of many varieties of oats, a number of grasses and three varieties of barley became infected. Flecking was observed on Lolium italicum (but not on L. perenne), and also on rye and wheat, but no pustules were formed on these plants.

A brief account of the histology of this rust was given by Pole-Evans (17). Ruttle and Fraser (18) published a detailed account of the cytology of Puccinia coronata on Banner oats, a susceptible variety, and on Cowra 35, a variety which showed different degrees of resistance.

Establishment of the uredospore culture

Grey Winter oats were used as the host for the greenhouse cultivation of the rust. The original source of the rust was from infected oats found at the University Farm, Cambridge, on November 14, 1927. The culture was maintained in the greenhouse for two years.

PART A. INOCULATION WITH UREDOSPORES OF PUCCINIA CORONATA

On May 29, 1928, seedlings of the following plants were inoculated with uredospores: Grey Winter oats, Wilhelmina wheat, Spratt Archer barley, rye, Lolium perenne and L. italicum.

Marked leaf areas from all these plants were removed one, two, three, five, seven, ten and fourteen days, respectively, after inocu-

lation, and prepared for microscopic study.

Macroscopic observations of the inoculated plants

Flecks appeared only on oats and wheat on the fourth day. By the sixth day barley showed signs of flecking, but these were faint, compared with those on oats and wheat, those on wheat being most distinct. By the tenth day oats, wheat and barley were definitely flecked, but there were no signs of flecking on the other plants. On the fourteenth day, when the final fixations were made, pustules were numerous on oats.

Inoculated leaf areas, which had not been removed during the course of the experiment, were kept under observation for one month. No pustules developed on wheat or barley, and there was no sign of infection on rye, *Lolium perenne*, or *L. italicum*.

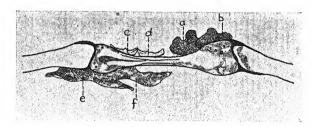


Fig. 10. $(\times 630.)$

Development of Puccinia coronata in oats (from uredospores)

The spores germinated by the end of the first day and formed appressoria over the stomata. The appressoria were either regular in outline or lobed as described by Ruttle and Fraser (18). Stomata of entry were not damaged. The appressorial contents had mostly passed into vesicles in the substomatal chambers, and by the end of the first day these had already assumed the elongate form characteristic of this rust. It was not unusual to find two or more vesicles in one substomatal cavity, as, for example, in Fig. 10 (e, f) (twenty-four-hour material), which shows four appressoria.

The rust made little progress during the next four days, though sometimes long, delicate infecting hyphae developed from one or both arms of the vesicles, leaving the latter empty. A median septum could often be seen in the vesicle. The infecting hyphae sometimes penetrated for some distance into the host tissue before forming a haustorium, though often the first host cell encountered was invaded.

At the end of the first week no extensive mycelium had developed, only a few delicate hyphae being observed among the mesophyll cells. By the tenth day, however, the stages preparatory to pustule formation were observed. Hyphae, dense with contents, were now massed

below the upper and lower epidermis, the hyphae in the central parts of the leaves being almost empty. Haustoria in these regions were drained; they were seen in both mesophyll and epidermal cells. By the fourteenth day numerous sporing pustules had formed on both surfaces of the leaves.

Throughout the development of the rust its relation with the host was completely congenial. The rate of development of *P. coronata* in Grey Winter oats in this experiment was considerably slower than its development in Banner oats described by Ruttle and Fraser (18).

Development of Puccinia coronata in wheat, rye, barley, Lolium perenne and Lolium italicum

The stage of development reached by the rust in all these plants

was approximately the same.

By the end of the first day many spores had germinated, and formed numerous appressoria over the stomata; the appressoria were usually

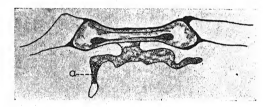


Fig. 11. (×630.)

empty, having formed large vesicles in the substomatal cavities. The vesicles were of the typical elongate form as found in oats, and infecting hyphae (a) had already begun to develop in many entries. This condition is shown in Fig. 11 (for rye).

Similar entries were also common in barley, Lolium perenne and L. italicum fixed at the same time. The stomata entered by the fungus

were not damaged at this stage.

After entry, the fungus made little progress. Later fixations showed that the vesicles with their short infecting hyphae had shrivelled in the substomatal cavities before reaching the mesophyll. This was common in all leaf areas fixed after the fifth day, and the guard cells of the stomata of entry had been killed. Only occasionally in these plants did infecting hyphae reach the mesophyll, when a few cells were either killed or their walls were thickened if in contact with the hyphae.

This is shown in Fig. 12, drawn from wheat fixed on the seventh day. Three germ tubes reached the stoma, but only one of these, a, entered, leaving outside the remnants of the appressorium. The other

two appressoria, b and c, appear to be drying up. The vesicle which developed from a produced two infecting hyphae, d and e, but no haustoria were formed. Hypha e put out several branches which passed between the mesophyll cells. No cells were killed, but their walls were often thickened where in contact with the hyphae. The fungus itself was somewhat shrivelled.

Only a single haustorium was observed in these plants. This was found in a leaf area of barley fixed on the seventh day. This entry was the furthest development of *Puccinia coronata* in barley, and in forming one small haustorium, it exceeded the progress of the rust in wheat, rye, *Lolium perenne* and *L. italicum*. The damage done to the leaf tissue was always negligible.

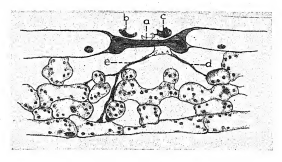


Fig. 12. (×315.)

Discussion of results with uredospores of Puccinia coronata Corda

The microscopic study of this rust in oats showed that in its normal development there was a congenial relation between host and parasite.

The rust gained entry into wheat, barley, rye, Lolium perenne and L. italicum, without at first damaging the stomata. Its subsequent progress was extremely limited, and only once did an infecting hypha produce a haustorium. This occurred in barley. Sometimes a few dead mesophyll cells were seen in contact with the infecting hyphae. Damage done to these plants was negligible and they were immune to attack from Puccinia coronata Corda.

PART B. INOCULATIONS WITH AECIDIOSPORES FROM RHAMNUS CATHARTICUS AND RHAMNUS FRANGULA

In the experiments with aecidiospores from R. catharticus and R. Frangula the method of inoculation was the same as in the previous experiments.

Infected bushes of both species of *Rhamnus* were found near Cambridge. Fresh spores were used to inoculate Grey Winter oats, Wilhelmina wheat, Spratt Archer barley, rye (unknown variety), *Lolium perenne* and *L. italicum*.

For each experiment a fresh supply of aecidiospores was obtained from the field. The spores invariably showed 100 per cent. germina-

tion in tap water when used for inoculation.

The experiments are summarised in Table V.

Table V. Experiments with aecidiospores from Rhamnus

Exp. No.	Date	Plants inoculated	Source of aecidiospores	Results
I	June 11, 1929	Oats Wheat Barley Rye L. perenne L. italicum	R. catharticus	Faint flecks on wheat and rye 6 days after inoculation. After 30 days no sign of infection of the other plants and no further development in wheat and rye; no sign of pustules
2	June 18, 1929	As above	R. catharticus	As in Exp. No. 1
3	July 4, 1929	As above	R. Frangula	No flecking on any of the plants. No sign of infection at the end of 30 days
4	July 17, 1929	As above	R. catharticus	Faint flecks on wheat on the 6th day and on oats on the 7th. One month after inoculation some leaves of oats and wheat still showed flecking but no pustules. No sign of infection on the other plants

In Exps. Nos. 1 and 2 marked leaf areas were removed at intervals from all the inoculated plants and prepared for microscopic study. The times of fixation were as follows: in the former one, three, seven and fifteen days after inoculation; in the latter two, three, seven, ten and fifteen days after inoculation.

The results of these experiments were unexpected, in that no pustules were produced on oats by aecidiospores from *Rhamnus catharticus*. Further consideration of these observations will be postponed until the discussion at the end of this section.

Microscopic observations on plants inoculated with aecidiospores from *Rhamnus catharticus*

Examination of the inoculated leaf areas showed that entries resulting from germinating aecidiospores are essentially the same as those resulting from the uredospores of *Puccinia coronata* Corda, described above. So far as is known, this study constitutes the first detailed demonstration of the method by which aecidiospores effect entry into the plant.

Although many spores germinated during the first twenty-four hours, germinating spores were observed on leaves as late as two

weeks after inoculation. The fungi entered the leaves of all plants inoculated. Appressoria were formed over the stomata; these were always smooth in outline, and never lobed as were often the appressoria which developed from uredospores of *P. coronata*. The vesicles, however, were almost identical with those which developed from the uredospores. Although the progress made after entry was slight, it varied in the different plants which were inoculated.

Development of the fungus in oats

Many aecidiospores germinated during the first twenty-four hours and formed appressoria over the stomata. Vesicles were found in the substomatal cavities, but no infecting hyphae had developed. The stomata of entry were not damaged.

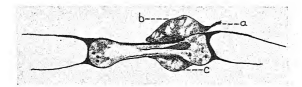


Fig. 13. $(\times 630.)$

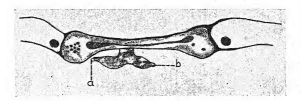


Fig. 14. (×630.)

The method of entry is shown in Figs. 13 and 14, drawn from twenty-four-hour material. In Fig. 13 a vesicle (c) is just being formed, the large appressorium (b) over the stoma still retaining most of its contents. Part of the shrivelled germ tube (a) is still attached to the appressorium.

The vesicle shown in Fig. 14 is fully developed and has the characteristic shape of the vesicle of crown rust of oats. Rudimentary infecting hyphae (a, b) can be seen, one at each arm of the vesicle.

In five-day material the fungus was still at the same stage, although the vesicles generally showed signs of shrivelling. Occasionally infecting hyphae had developed and had reached mesophyll cells, but the vesicles were always empty and shrunken and the infecting hyphae considerably shrivelled. No haustoria had developed. One of these entries is shown in Fig. 15. Here the guard cells are still living but are somewhat discoloured. The empty germ tube (a) and appressorium (b) are seen outside and the shrunken vesicle (c) in the substomatal chamber. Only one infecting hypha (d) has developed. This reached a mesophyll cell (e), which is dead and deeply stained. It is impossible to distinguish the contents, and therefore it is not known whether a haustorium was formed.

Similar entries were found in leaf areas fixed on the tenth and fifteenth day after inoculation, but no progress beyond the stage shown in Fig. 15 had been made. In all the leaf areas of oats not a single haustorium was seen and there was no trace of an inter-

cellular mycelium.

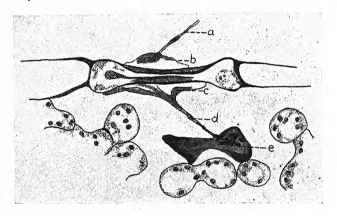


Fig. 15. (×630.)

Development of the fungus in Lolium perenne and Lolium italicum

The spores germinated, forming appressoria and vesicles. These were numerous in both species of *Lolium* by the end of the first day and were similar to those described for oats. No further development occurred after entry. Leaf areas fixed on the fifteenth day showed empty, shrivelled vesicles but no infecting hyphae, although rudiments of these were sometimes apparent.

Development of the fungus in wheat

Vesicles were quite numerous in wheat by the end of the first day. Frequently two or more were found in the same cavity, and no damage to the stomata resulted. No infecting hyphae developed beyond the rudimentary stage, as in *Lolium perenne* and *L. italicum*.

Inoculated areas of wheat which showed flecking at the time of fixation revealed no development of the fungus after entry. The

Results of Inoculating Cereals with Spores of Cereal Rusts 281

formation of vesicles marked the maximum development in wheat. Vesicles were found on the fifteenth day after inoculation, but at this time they were always shrivelled.

Development of the fungus in barley

Leaf areas of barley showed numerous entries; frequently as many as four vesicles were found in one substomatal cavity.

An unusual entry found in barley is drawn in Fig. 16 (five-day material). The remains of two appressoria (a, b) are seen over the stoma. Separate entries were made, but these appear to have fused

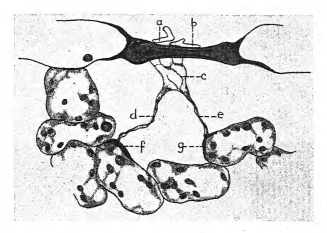


Fig. 16. (×630.)

to form a large single vesicle (e) in the cavity. This is now empty and somewhat shrivelled. Two infecting hyphae (d, e) were produced, both of which reached the mesophyll but did not form haustoria. The walls of the mesophyll cells are thickened (f, g) where they are in contact with the hyphae, but both cells are living.

The furthest development of the fungus in barley was seen in a five-day fixation in which an infecting hypha formed a large haustorium in a mesophyll cell. No further development took place. This was the only haustorium seen in barley. There was no trace of an intercellular mycelium in barley.

Vesicles were found as late as the fifteenth day, the guard cells of the stomata of entry being usually dead. The vesicles at this time were either empty or completely shrivelled.

Development of the fungus in rye

Entries were common in rye by the end of the first day; these were similar to those described in oats. As late as the fifteenth day,

however, germinating spores were still seen on the leaves.

Rye only, of all the plants inoculated with aecidiospores, showed traces of an intercellular mycelium. This was found in several areas fixed on the fifteenth day. The mycelium was scanty and consisted of only a few meagre hyphae. There were no signs of the entries from which the mycelia had developed, except that collapsed stomata were found in the infected regions. Large haustoria were seen in some cells, the latter being usually empty. These haustoria were of distinctive

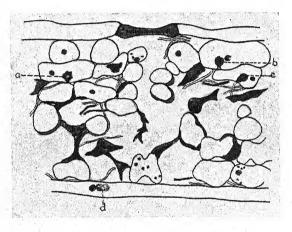


Fig. 17. (×315.)

type with thick necks and enclosed in striated sheaths of irregular margin. Some haustoria were transparent, with deeply stained sheaths, while others were deeply stained and the sheaths were clear, colourless and plainly striated. Haustoria similar to these found in rye have been described by Allen (2), occurring in infections of Puccinia graminis Tritici on Mindum wheat, a resistant variety.

Fig. 17 (fifteen-day material) is drawn from a section through one of these infected regions in rye in which an intercellular mycelium had developed. Most of the cells are dead and empty; in some the cell walls are very much thickened, while other cells have collapsed and are deeply stained. The hyphae and haustoria are dead. Haustoria are present in the mesophyll cells at a, b and c. These are stained in different ways and show the characteristic thick necks. Only haustorium d, which is in contact with the host nucleus, appears to

Discussion of inoculations with Aecidiospores from *Rhamnus* spp.

The three experiments outlined in Table V show that the aecidiospores from R. catharticus failed to produce pustules on any of the plants inoculated, although flecking occurred on wheat and rye in two of the experiments (Nos. 1 and 2), and on wheat, rye and oats in the other experiment (No. 4). In the experiment (No. 3) with aecidiospores from R. Frangula there were no signs of infection on any of the inoculated plants.

The microscopic study, which was restricted to plants inoculated with aecidiospores from R. catharticus, has shown that the failure of the fungus to produce successful infection was not due to failure to

make entry, since entries were numerous in all these plants.

Although the appressoria formed by the germinating aecidiospores were always smooth in outline, and never lobed as were some appressoria from germinating uredospores of crown rust of oats, the vesicles which developed had the characteristic form peculiar to *Puccinia coronata* Corda.

The microscopic study showed, however, that no true infection of oats resulted from inoculation with aecidiospores from *Rhamnus catharticus*. Entries were made and a few infecting hyphae developed, but no haustoria were formed.

In wheat, Lolium perenne and L. italicum the progress of the fungus was similarly restricted, no more development being made than in oats. It seems, therefore, that the flecks seen on wheat and oats were

not due to the fungus.

Although barley showed no flecking, the fungus made slightly more progress in this plant than in oats and wheat. The development of infecting hyphae was rather more frequent, and in one entry a haustorium was formed in a mesophyll cell. There was, however, no further development.

Only in rye did the fungus develop so far as to produce an intercellular mycelium. The mycelium was scanty and it was always clear that there was an antagonistic relation between rye and the fungus, since the hyphae and most of the haustoria were dead, as well as

many of the mesophyll cells in the infected zones.

All the plants inoculated were immune to aecidiospores obtained from *Rhamnus catharticus*. It is evident, therefore, that the rust was not the aecidial stage of the form of *Puccinia coronata* Corda, which was cultivated on Grey Winter oats in the greenhouse. It doubtless belonged to one of the many other physiologic forms of *P. coronata*.

Section V. Puccinia graminis Secalis Erikss.

P. graminis Secalis is one of six specialised forms of black rust recognised by Eriksson (7). According to him, this rust attacks rye, barley, Agropyron repens (couch-grass), and other grasses in Sweden, but not wheat or oats.

Stakman and Piemeisel (23) found that *Puccinia graminis Secalis* from *Agropyron repens* infected rye, barley, *A. repens*, and some other grasses, but oats and wheat were rarely infected. Mehta (15) was able to infect rye, barley, *A. repens*, and one variety of wheat (Red Sudan),

but not oats.

Microscopic studies of *Puccinia graminis* have been confined almost exclusively to *P. graminis Tritici*, the black rust of wheat. A short account of the histology of *P. graminis Tritici* was published by Pole-Evans (17). Stakman (19) studied the histology of this rust on a susceptible variety of wheat (Minnesota No. 163) and on a resistant variety (Khapli). Further, he (20) made microscopic studies of oats inoculated with *P. graminis Tritici*, wheat inoculated with *P. graminis Avenae*, oats inoculated with *P. graminis Hordei*, and rye, wheat and barley inoculated with *P. graminis* from *Dactylis glomerata*.

There remain to be mentioned in connection with *Puccinia graminis* Tritici the study by Newton (16) of the development of the rust in a susceptible and a resistant variety of wheat, and the cytological

studies by Allen (1, 2, 3).

ESTABLISHMENT OF THE CULTURE ON AGROPYRON REPENS

Uredospores were collected at Cambridge and greenhouse cultures

were established on rye and couch-grass.

For some experiments uredospores for inoculation were taken directly from couch-grass out of doors, but for the later experiments they were obtained from the greenhouse cultures.

EXPERIMENTS WITH PUCCINIA GRAMINIS SECALIS

The plants inoculated with this form of black rust were as follows: Agropyron repens, rye (unknown variety), Spratt Archer barley, Wil-

helmina and Norka wheats, and Grey Winter oats.

The experiments with *Puccinia graminis Secalis* are given in Table VI. In Exp. 4 marked leaf areas were removed at intervals and prepared for microscopic study; the fixations were made one, three, five, seven and ten days after inoculation. In the remaining six experiments the plants were kept under observation for one month.

The data in the last column of the table may be summarised as

follows:

Couch-grass. In five out of the seven experiments medium to heavy infections resulted; in one experiment there was a weak infection, and in one experiment no infection. Out of a total of sixty-nine plants inoculated (omitting Exp. No. 4) thirty-nine produced pustules.

Table VI. Experiments with Puccinia graminis Secalis

Exp. No.	Date	Plants inoculated	Gulture used	Results
I	June 6, 1928	Couch-grass Rye Barley Wil. wheat Oats	From couch- grass in open	Couch-grass 9/12,* fair infection Rye 8/10, fair infection No sign of infection on: Barley 0/10 Wheat 0/10 Oats 0/10
2	Oct. 3, 1928	As in 1	From couch- grass in open	Couch-grass o/17, flecking Rye 6/12, weak infection Barley o/15, flecking Wheat o/14 Oats o/12
3	July 23, 1929	As in 1	G I from rye (greenhouse culture)	Couch-grass 10/10, very good infection Rye 10/10, very good infection Barley 0/10 Wheat 0/10 Rye 0/10
4	Aug. 25, 1929	Couch-grass Rye Barley Wil. wheat Norka wheat Oats	G 2 from couch-grass (greenhouse culture)	Areas removed from plants and fixed. There were numerous pustules on rye on 7th day and on couch-grass on 10th. Weak infection on barley after 21 days. No sign of infection on Wil. wheat, Norka wheat, or oats
5	Aug. 25, 1929	As in 4	G 3 from rye (greenhouse culture)	Couch-grass 10/10, excellent infection Rye 10/10, excellent infection Barley 2/10, very weak (after 21 days) Wil. wheat 1/10, very weak Norka wheat 0/10 Oats 0/10
6	Sept. 6, 1929	Couch-grass Rye Barley Wil. wheat Oats	G 4 from rye (greenhouse culture)	Couch-grass 8/10, good infection Rye 11/11, excellent infection Barley 0/10 Wil. wheat 0/9 Oats 0/7
7	Sept. 29, 1929	Couch-grass Rye Barley Wil. wheat Oats	G 5 from rye (greenhouse culture)	Couch-grass 2/10, weak infection Rye 3/10, weak infection Barley 0/10 Wil. wheat 0/10 Oats 0/10

^{*} The denominator of the fraction represents the number of plants inoculated, the numerator the number which produced pustules.

Rye. Medium to heavy infections resulted in five out of the seven experiments, and weak infections in the other two. Out of a total of sixty-three plants inoculated (omitting Exp. No. 4) forty-eight produced pustules.

Barley. Very weak infections on barley were observed in two experiments (Nos. 4 and 5), and flecking in one experiment (No. 2). There was no sign of infection in the other experiments. Out of a total of sixty-five plants inoculated (omitting Exp. No. 4) only two produced pustules.

Wilhelmina wheat. Out of a total of seventy-three plants inoculated in the seven experiments, only a single plant (in Exp. No. 5) produced pustules, and these were few and minute. Norka wheat and oats

were consistently immune.

These results are in agreement with the observations of Mehta (15), who found that couch-grass and rye were favourable hosts for *P. graminis Secalis*, and that Spratt Archer barley showed only weak infections. Mehta, and other workers, however, obtained good infections on other varieties of barley with *P. graminis Secalis*. Mehta also obtained weak infections on one variety of wheat, as was observed in my experiments with one plant of Wilhelmina wheat. Various workers have found that oats is consistently immune to this form of black rust.

Microscopic observations of plants inoculated with Puccinia graminis Secalis

Development of Puccinia graminis Secalis in Agropyron repens

Under the conditions of the experiment the rust made little progress in couch-grass during the first five days. By the end of the first day numerous appressoria had been formed over the stomata but only about I per cent. had formed vesicles and no infecting hyphae were seen at this time. Even on the third and fifth days comparatively few entries were found although appressoria were still numerous over the stomata. Usually, whether these stomata were penetrated or not, the guard cells were dead, and had lost their affinity for the diamant fuchsin stain. Stomata similar to these were also found by Allen (5, 3) in infections of wheat with *P. graminis Tritici*.

Sometimes guard cells were seen in which part of the protoplast was normally stained and part seemed dead and shrunken. This condition was seen only sometimes where the fungus had entered, as shown for example in Fig. 18 (seven-day material). The shrivelled germ tube (a) and appressorium (b) are seen outside. The fungus entered at one end of the stoma, forming a large vesicle (c). A single infecting hypha (d) was produced which ran more or less parallel to the leaf surface, and gave rise to a large haustorium (e) in the epidermal cell adjacent to the stoma of entry. The part of the guard cell nearest the haustorium is dead and shrunken (f), whereas the end where the fungus entered is apparently undamaged. The formation

of the first haustorium in an epidermal cell as in this entry was found

to be general in infections with P. graminis Secalis.

From the seventh to the tenth day all stages of reproductive activity could be seen. When the final fixations were made (ten days after inoculation) numerous pustules had been formed on both surfaces of the leaves. Most of the intercellular hyphae in the pustule regions were quite empty, and were frequently septate. They were so densely packed that the outlines of some of the cells were completely obscured. Haustoria were rare in host cells immediately below pustules, and when present they were drained.

Apart from the death of guard cells, the relation between couch-

grass and the rust throughout its development was congenial.

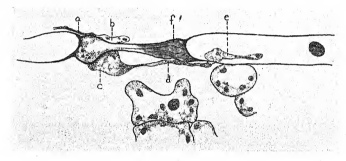


Fig. 18. (×630.)

Development of Puccinia graminis Secalis in rye

In rye numerous appressoria were seen over the stomata twenty-four hours after inoculation, and although few entries had been made, they exceeded in number those found in couch-grass at the same time. The guard cells of the stomata of entry were usually dead and discoloured as in couch-grass. Approximately 5 per cent. of the appressoria had produced vesicles in rye by the end of the first day, but no infecting hyphae had been formed.

The fungus advanced more rapidly in rye than in couch-grass. By the third day long infecting hyphae had developed, and many of these had produced haustoria. The infecting hyphae always took a course parallel to the leaf surface, and the first haustorium was formed in the epidermal cell adjacent to the stoma of entry.

By the fifth day an intercellular mycelium was well developed, and haustoria were numerous in both mesophyll and epidermal cells. Leaf areas fixed at this time showed that vesicles were still being formed in the substomatal cavities.

On the seventh day numerous pustules were found on both surfaces

of the leaves. The mycelium was dense in the mesophyll tissue underlying the pustules.

As in couch-grass, the relation between host and rust was con-

genial.

Development of Puccinia graminis Secalis in barley

The early development of the rust in barley was similar to that in rye. Appressoria were abundant by the end of the first day, but the fungus had rarely entered the leaves. The fungus had already shrivelled in one or two of the entries.

New vesicles were still being formed on the third day. No haustoria were seen in barley until the fifth day, and they were comparatively rare. As always in this rust, the first haustorium was formed in an epidermal cell adjacent to the stoma of entry. Haustoria were

smaller than those in couch-grass and rye.

Leaf areas fixed seven and ten days after inoculation still showed no trace of an intercellular mycelium. Appressoria were still seen over the stomata, and occasionally empty vesicles were found in the substomatal cavities. The guard cells of the entered stomata were invariably dead and discoloured. It was apparent that barley was not a favourable host for *P. graminis Secalis* under the conditions of the experiment.

No material was fixed after the tenth day. In this particular experiment (No. 4, Table VI), a few minute pustules appeared on some leaves twenty-one days after inoculation. It is evident therefore that some entries resulted in the establishment of an intercellular mycelium, but since no material was fixed nothing can be said concerning the host-parasite relation in these infections which produced minute

pustules.

Development of Puccinia graminis Secalis in wheat

In Wilhelmina wheat, only one small vesicle was seen in the leaf areas fixed twenty-four hours after inoculation, although many appressoria were seen over the stomata. The guard cells were always dead.

The rust made little progress during the next two days, and entries were rare. On the fifth day vesicles were more common, but they were usually empty. Some had given rise to infecting hyphae but these were poorly developed and none had formed haustoria.

The seven- and ten-day fixations showed no further development of the fungus. Not a single haustorium was found in any of the leaf

areas.

Apart from the dead guard cells where the fungus was seen, there was no damage to the leaf tissues, the fungus making no progress after entry.

The development of the rust in Norka wheat was similar to that in Wilhelmina wheat. A few entries were seen, but no haustoria were formed.

In this particular experiment (No. 4, Table VI) neither variety of wheat became infected, but it should be pointed out that in one of the other experiments (No. 6, Table VI) a weak infection was obtained on Wilhelmina wheat.

Development of Puccinia graminis Secalis in oats

Appressoria were formed over the stomata, but few entries were made in oats. The guard cells of stomata bearing appressoria were always dead, being discoloured whether the fungus entered or not.

A few vesicles were found twenty-four hours after inoculation, but no infecting hyphae had developed. Newly formed vesicles were seen in the later fixations, occasionally even as late as the tenth day, but neither infecting hyphae nor haustoria were seen in oats.

Apart from the killed guard cells there was no damage to the leaf tissues in oats.

DISCUSSION OF EXPERIMENTS WITH PUCCINIA GRAMINIS SECALIS

As was pointed out earlier, macroscopic observations of the experiments with P. graminis Secalis indicate that only rye and couch-grass, amongst the plants inoculated, were susceptible to this rust. Two plants of barley, out of a total of sixty-five inoculated, produced pustules, and these were minute and late in developing. Similarly, in Wilhelmina wheat, a weak infection resulted on one plant, out of a total of seventy-three inoculated. Norka wheat and oats were consistently immune.

Microscopic study of fixed leaf areas from Exp. 4 showed that the

fungus entered all these plants.

Rye and couch-grass were the only true hosts for the rust. From the beginning the fungus developed more rapidly in rye, and pustules

were formed three days earlier than in couch-grass.

In barley infecting hyphae occasionally produced haustoria in the epidermal cells adjacent to the stomata of entry, but no further progress was made. In wheat no further development beyond the formation of infecting hyphae was made, while in oats no infecting

hyphae developed from the vesicles.

In all the plants inoculated it was found that, with very few exceptions, the guard cells of the stomata bearing appressoria were killed, whether or not entry was made. Sometimes, in entries in rye and couch-grass, it seemed that only part of the guard cell was injured, and it is perhaps significant that, in all such cases, the discoloured and shrunken end of the guard cell was found always to be the end which was in contact with the epidermal cell in which the first

haustorium was produced.

Finally, mention should be made of the interesting peculiarity that in *P. graminis Secalis* the first haustorium is always formed in an epidermal cell adjacent to the stoma of entry.

SUMMARY

1. Inoculation experiments have been carried out with uredospores of *Puccinia triticina* Erikss., *P. glumarum Tritici* Erikss., *P. anomala* Rostrup, *P. coronata* Corda, *P. graminis Secalis* Erikss., both on their normal hosts and on cereals on which they do not normally occur ("inappropriate hosts").

2. The development of these fungi in the "inappropriate hosts" has been compared with their development in their normal hosts.

3. In general, the relation of the host to the fungus which does not develop on it in nature is one of antagonism, for the mesophyll cells near the stomata of entry are usually killed. On the other hand, the initiation of invasion by these fungi on the "wrong hosts" proceeds normally at first. Exceptionally, the fungus on the "wrong host" kills the guard cells of the stomata of entry, but, failing to develop further than the formation of substomatal vesicles, leads to no further antagonistic reaction on the part of the host.

4. P. triticina is not very sharply specialised to wheat, for it sometimes produces normal pustules of uredospores on rye. The different types of reaction between this rust and rye are described. Occasionally minute abortive pustules of this rust were produced on barley, but the relation is usually antagonistic. In oats progress by this rust

was negligible, and only once was a haustorium seen.

5. P. glumarum Tritici produced intercellular mycelia in rye and barley, but no pustules were formed. Although the fungus entered

oats no mycelium developed.

6. P. anomala initiated invasion of wheat, rye and oats, killing the guard cells of the stomata of entry but failing to form mycelia in the tissues.

7. P. coronata Corda (uredospores from oats) entered wheat, barley, rye, Lolium perenne and L. italicum, but usually failed to develop beyond the formation of infecting hyphae. The contiguous mesophyll cells were sometimes killed.

8. The initiation of infection by aecidiospores of *Puccinia coronata* Corda from *Rhamnus catharticus* is described on oats, wheat, rye, barley and *Lolium* spp. Progress of the fungus in these hosts was negligible (except in rye), and no uredospores were formed. It is concluded that this biologic form has a host range outside the hosts experimented with.

Results of Inoculating Cereals with Spores of Cereal Rusts 291

o. Puccinia graminis Secalis from rye and Agropyron repens entered wheat, barley and oats, and occasionally produced small pustules of uredospores on barley and wheat. No infecting hyphae were produced from the substomatal vesicles in oats. In the development of Puccinia graminis Secalis the first haustorium is always formed in an epidermal cell adjacent to the stoma of entry.

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AN INVESTIGATION OF THE TEMPERATURES LETHAL TO SOME WOOD-DECAYING FUNGI

By H. B. S. MONTGOMERY, B.A.

Introduction

Timber which is apparently sound and serviceable may contain traces of infection which become obvious only when the timber is incorporated in structures and decay has advanced considerably. It is desirable to heat timber suspected of being thus infected in order to kill any fungi present. It sometimes happens that valuable ornamental carved timber becomes infected with a wood-decaying fungus, and the question arises whether it is possible to kill the fungus without damaging the wood. There is thus a need for accurate data concerning the time required at certain temperatures to kill various wood-decaying fungi under known conditions (age, moisture content,

etc.).

The literature on the subject is very limited. Snell (3) exposed wood in various stages of decay to heat in sealed Mason jars. He used both dry and moist wood, and showed that a fungus in dry wood resists heat treatment much better than a fungus in moist wood. He worked with rather long time periods, combined with fairly low temperatures. Hubert (1) subjected blocks of various sizes of naturally infected timber to heat for different time periods, chiefly in a Tiemann dry kiln, to determine whether commercial kiln processes were efficient in killing wood-decaying and sap-staining fungi in timber. His time intervals were wide and his results show no great differences in resistance among the various wood-decaying fungi tested. Liese (5) experimented with agar cultures of the fungus and short periods of time at fairly high temperatures.

The following experiments were undertaken to find out what relation, if any, exists between results obtained with agar cultures and those with infected wood blocks, and also to obtain some preliminary figures for the lowest temperatures which would be lethal to various

wood-decaying fungi in a reasonably short time.

Cultures of a number of common wood-decaying fungi were obtained from the Forest Products Research Laboratory, Princes Risborough, and subcultured on 2 per cent. malt agar. The fungi used were Merulius lacrymans (Wulf.) Fr., Poria vaporaria (Pers.) Fr., Pholiota adiposa Fr., Lentinus lepideus Fr., Lenzites abietina Bull., Lenzites saepiaria (Wulf.) Fr., L. striata Swartz, L. trabea Pers., Polystictus versicolor (Linn.)

Fr., Polyporus hispidus (Bull.) Fr., Schizophyllum commune Fr., and Fomes fraxineus (Bull.) Fr.

EXPERIMENTAL METHODS

For the purpose of these tests, the fungi were cultured in small blocks $3 \times 1 \times 1$ in. of ash or Scots' pine wood as follows:

ASH
Lenzites striata
Lenzites trabea
Polystictus versicolor
Polyporus hispidus
Schizophyllum commune
Fomes fraxineus

Scots' Pine Merulius lacrymans Poria vaporaria Pholiota adiposa Lenzites abietina Lenzites saepiaria Lentinus lepideus

After sterilisation by autoclaving, these blocks were inoculated by placing eight to ten of them on an active malt agar culture of each fungus in a suitable glass vessel. These blocks were allowed to decay for nine months before use, so that the fungus had thoroughly permeated the blocks.

The fungi were also cultured in $\frac{3}{4}$ in. test-tubes containing 10 c.c. (approx.) of 2 per cent. malt agar as slants, and the heat treatments were carried out on these cultures when they were one month old. The treatment was applied by immersing a number of these test-tube cultures in a large bath of water maintained at the desired temperature in a thermostatically controlled oven. At each time interval, one tube culture was removed and allowed to cool. Ten pieces of the culture were then removed from the centre of the slant and transplanted on to a plate of 2 per cent. malt agar. The plates were incubated at a temperature suitable to the fungus (generally 20° C.) for at least one month, and a note was made of any growth which occurred. Where no growth occurred from any of the inocula in a plate, it was concluded that the heat treatment had killed the fungus. The results of these tests are presented in Table I.

It is most important that the inocula should be taken from a standard position in the slant. Inocula derived from the upper end of the slant proved more resistant to heat than inocula from the centre. This resistance seems to be due, in part, to the less moist

conditions prevailing at the upper end.

Using these preliminary data as a guide, tests were made with the infected wood material. The $3 \times 1 \times 1$ in. blocks were cut into 1 in. cubes, and all the cubes of wood infected with any one fungus were stored in a glass dish and kept moist. By this means it was hoped to equalise the moisture contents of the blocks for each fungus. The blocks were kept in this manner for periods of over three weeks and up to two months. This proved very satisfactory from the point of view of moisture content, but it permitted moulds to grow on the surface of the blocks. These were troublesome only on wood infected

Table I. Results with fungi in agar

		4	o° G.	2000			in agan ° C.		6-9-0					
							<u> </u>		60° C.					
	15 min.	30 min	. 45 . min	. mir		30 . min.	60 min.	120 min.	15 min.	30 min.	60 min.			
Merulius lacrymans	•	•			•									
	•	•	•	•	•	•	•							
	•	•	•	•	•		•		_		• 3			
	<u>.</u>	-		<u>:</u>	_	_	_		•	•	•			
Poria vaporaria				_	•	•	•	•	•	•	• .			
•				•	•	•	•	•		•	_			
		•				•	•		_	_				
						_	-	<u> </u>						
Lenzites abietina														
	•	•	•						•					
	•	•							_	_	-			
	•	•	•	•										
Pholiota adiposa	•	• •	:	•	_		_	_						
notivia aarposa	•	•	•	•	•	•	•	•		-	_			
	•	•	•	•	•	•	•	•	_	_				
Polyporus hispidus	•	•	•	•	_	-	_	-	•		•			
copporus inspinus	•	• ,	•	•	•	•	•	•		_	_			
	•	•	•	•	•	•	•	•	4000	_				
	i	i	+	i	_	_	_		•		•			
Polystictus versicolor	-1-	T	Т	Т	•	•	• •	•	•	•	•			
	·	•	•	•	•	•	•	•	•	•	_			
		:		• :	•	•	•	•	•	_	_			
					<u>;</u>	÷	·	<u>.</u>	<u>.</u>	_				
	+	+	+	+		,								
				·		60° C		•	·	Α.				
	30		6o											
	30													
	min.	45 min.	min. n	75 9	00 120 in. min.		80 240 in mir	300 300 min.	360 42 min. m	20 480 in. min	540 . min			
Lenzites trabea	min.	min.	min. n		in. min.	min. n		300 n. min.	360 42 min. m					
	min.	min.	min. n		in. min. . – + +	150 I min. n		300 n. min.	360 42 min. m					
	min.	min.	min. n + + +		in. min.	150 I min. n	in. mir – –	300 n. min.	360 42 min. m					
Lenzites trabea	min.	min.	min. n + + + + +	:	in. min.	150 I min. n	in. mir – –	9 300 n. min.	360 42 min. m					
Lenzites trabea	min. : +	min.	min. n + + + + +	: +	in. min.	150 I min. n	in. mir – –	300 300 min.	360 42 min. m					
Lenzites trabea	min. : +	min. : : : +	min. n + + + + + + +	: + •	in. min.	150 I min. n	in. mir – –	300 n. min.	360 45 min. m					
Lenzites trabea	min.	min.	min. n + + + + +	: +	in. min.	150 I min. n	in. mir – –	300 300 min.	360 42 min. m					
Lenzites trabea	min. : +	min. : : : + +	min. n + + + + + - + + -	· + · · + · · · · · · · · · · · · · · ·	nin. min	150 I min. n	in. mir – –	9 300 n. min.	360 42 min. m					
Lenzites trabea L. saepiaria	min. : +	min. : : : + + -	min. n + + + + + - + + + + - +	: + : + - - +	in. min		nin. mir	9 300 n. min.	360 42 min. m					
Lenzites trabea L. saepiaria	min. + + + + + +	min.	min. n + + + + + + + + + + + + + +	·	nin. min		nin. mir	9 300 n. min.	360 42 min. m					
Lenzites trabea L. saepiaria	min. : +	min. : : : + + -	min. n + + + + + + + + + + + + + + + + + +	: + : + - - +	in. min		nin. mir	9 300 n. min.	min. m	in. min				
Lenzites trabea L. saepiaria	min. + + + + + +	min.	min. n + + + + + + + + + + + + + +	· · · · · · · · · · · · · · · · · · ·	in. min		nin. mir	n. min.	min. m	in. min				
Lenzites trabea L. saepiaria Lentinus lepideus	min. + + + + + +	min.	min. n + + + + + + + + + + + + + + + + + + +	· · · · · · · · · · · · · · · · · · ·	nin. min		nin. mir	0 300 min	min. m	in. min				
Lenzites trabea L. saepiaria Lentinus lepideus Schizophyllum	min. + + + + + +	min.	min. n + + + + + + + + + + + + + + + + + +	· · · · · · · · · · · · · · · · · · ·	in. min		nin. mir	n. min.	min. m	in, min				
Lenzites trabea L. saepiaria Lentinus lepideus	min. + + + + + +	min.	min. n + + + + + + + + + + + + + + + + + + +	· · · · · · · · · · · · · · · · · · ·	nin. min		nin. mir	n. min.	min. m	in. min. min				
Lenzites trabea L. saepiaria Lentinus lepideus Schizophyllum	min. + + + + + +	min.	min. n + + + + + + + + + + + + + + + + + + +	· · · · · · · · · · · · · · · · · · ·	nin. min		nin. mir	n. min.	min. m	in. min. min				
Lenzites trabea L. saepiaria Lentinus lepideus Schizophyllum commune	min. + + + + + +	min.	min. n + + + + + + + + + + + + + + + + + + +	· · · · · · · · · · · · · · · · · · ·	nin. min		nin. mir	n. min.	min. m	in. min				
Lenzites trabea L. saepiaria Lentinus lepideus Schizophyllum commune	min. + + + + + +	min.	min. n + + + + + + + + - + + - + + - + + - + + - + + - + + +	· · · · · · · · · · · · · · · · · · ·	nin. min		nin. mir	n. min.	min. m	in. min. min				
Lenzites trabea L. saepiaria Lentinus lepideus Schizophyllum commune	min. + + + + + +	min.	min. n + + + + + + + + + + + + + + + + + + +		nin. min		nin. mir	n. min.	min. m	in. min. min				
Lenzites trabea L. saepiaria Lentinus lepideus Schizophyllum commune	min. + + + + + +	min	min. n + + + + + + + + - + + - + + - + + - + + - + + - + + +		nin. min		nin. mir	n. min.	min. m	in. min. min				
Lenzites trabea L. saepiaria Lentinus lepideus Schizophyllum commune	min. + + + + + +	min	min. n + + + + + + + + + + + + + + + + + + +	· · · · · · · · · · · · · · · · · · ·	in. min		nin. mîr.	n. min.	min. m	in. min. min				
Lenzites trabea L. saepiaria Lentinus lepideus Schizophyllum commune Lenzites striata	min. + + + + + +	min	min. n + + + + + + + + + + + + + + + + + + +	• • • • • • • • • • • • • • • • • • • •	in. min		nin. mir	n. min.	min. m	in. min. min				
Lenzites trabea L. saepiaria Lentinus lepideus Schizophyllum commune Lenzites striata	min. + + + + + +	min	min. n + + + + + + + + + + + + + + + + + + +	· · · · · · · · · · · · · · · · · · ·	in. min		nin. mîr.	n. min.	min. m	in. min. min				
Lenzites trabea L. saepiaria Lentinus lepideus Schizophyllum commune Lenzites striata	min. + + + + + +	min	min. n + + + + + + + + + + + + + + + + + + +	• • • • • • • • • • • • • • • • • • • •	in. min		nin. mîr.	n. min.	min. m	in. min. min				
Lenzites trabea L. saepiaria Lentinus lepideus Schizophyllum	min. + + + + + +	min	min. n + + + + + + + + + + + + + + + + + + +	• • • • • • • • • • • • • • • • • • • •	in. min.		nin. mîr.	n. min.	min. m	in. min. min				

Note. Each — or + indicates the absence or presence of fungal growth from a single test-tube culture.

with certain fungi, ash wood infected with Polyporus hispidus being

particularly prone to mould attack.

Various methods of heating the wood blocks in a water bath, avoiding loss or gain of moisture, were considered. It was thought undesirable to put them in a glass vessel—such as a test-tube or a Mason jar-for treatment, since heating to the desired temperature would be a slow process, and further, exchange of moisture might occur between the block and the relatively large volume of air (either saturated or dry) in the glass vessel. Experiments were then made with methods of coating the block with some impervious paint or varnish, but, apart from the risk of vapours from the paint having a toxic effect, it was found almost impossible to render the block watertight. Finally, thin sheet rubber was used. This was first cemented with rubber solution into tubes about 4 in. long and wide enough to accommodate conveniently a 1 in. cube. After putting a block into a tube, the ends were tightly tied with twine, thus forming a watertight bag. By this method, a very small volume of free air was included and the block was in close contact with the source of heat. The rubber bags, each containing one block, were submerged, by means of a wire basket, in a large bath of water maintained at the desired temperature.* One block was removed at the end of each time period, and, after cooling, it was flamed on the surface and split in half. From this freshly exposed surface of each half, five small splints were removed and placed on prune agar in a Petri dish. Thus, in all, ten splints of wood were removed from the centre of each block. The inoculated dishes were stored at 20° C. (approx.), and where no growth occurred after one month, the fungus was considered killed. The figures are presented in Table II. To keep a check on the moisture content of the blocks during treatment, a control block was enclosed in a rubber bag, and submerged in the bath for the course of the treatment, and its moisture content was determined after cooling as a percentage of oven-dry weight. This moisture content was assumed to be similar to that of the other blocks similarly decayed and heat treated.

In view of the uniformity of the results, it would seem that the variation in moisture content of these blocks was not large enough to affect the resistance of the fungus to heat. It is also of interest that these moisture contents all permitted the fungi to grow freely.

^{*} The temperature was kept constant by manipulation of a fine adjustment on the gas supply, a mechanical thermostat not being available. The temperature varied \pm 1° C.

Table II. Results with fungi in wood

			40°	C.			<i>J</i>	0						
	I 5 min.	30 min.	60 min.	90 min.	120 min.	150 min.	I mi	5 in.	30 min.		60 min	90 min	120 min	Average % moisture
Merulius lacrymans	_		_	_		•		•	•		•			103
	-	_	_		•	•			•	•	•			Ü
Poria vaporaria	+	+	+	+	+	•		•	•	•	•	•	•	59
		•	•	·	9		_	-	<u>.</u>	÷		<u>.</u>	÷	39
	•	•	•	•		•	-	-	_	_	_	-	_	
Lenzites abietina	•	•	+	+	+	•	-	_	_	-	_		-	105
			·			в •	-	-	<u>.</u>	÷	<u>:</u>	<u>.</u>	•	103
	•	•			•	•	-	-	_	_	_		_	
	•	•	•	•	•	•	-		_	_		_		
Pholiota adiposa	+	:	+	÷	+	<u>.</u>		-	-	-		-		35
•			•				-			_	_	_	_	33
	•	•	•	•	•	٠	-	_	_	-	-	-		
Polyporus hispidus	•	•	•	•	•	•		_	_	_	_	_	_	46
- opposite mapation		•	·	·	·	·	-	_	_		- - -			7.
	•	•	•		• •	•	-	_	_	_	_	•	•	
Polystictus versicolor	•	•	•	•	•	•	-	_		_		=	·	55
1 diysticius cersilotor	·	:	:	:	:			+ + +	++	+	+ - +	_	_	33
			•		•			÷	÷	+	+	-	_	
				. ~						•				
			60	°a.			_			65	°a.			_ Average
	nin.	go min.	60 min	90 min.	120 min	150 min.	'n	15 nin	. mi	o n. n	60 nin.	90 min.	120 min.	% moisture
Lenzites trabea				_	_	_								46
	- 1-	_		_	_	_		•			•	•	•	
	÷	++	++	+	_	-		•	•		•	•	•	
$L.\ saepiaria$	•		+	_	_	÷		:	:		:	÷		63
•		+	+	+	+	-		•			, ·	•		
	•	++++	+	_	-	_		•	•		•	•	•	
	•	+		_		_		:			:	÷		
Lentinus lepideus									4	-	_	_	_	45
			•					٠	4	-	_	_		
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Schizophyllum		•						•	-	-	-	•		31
commune	•	•	•	•	•	•		+	-	-	_ , .	•	_	
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Lenzites striata		:	•	- 25					· -	-	-		_	61
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Fomes fraxineus	•			-	-	:				_	_	_	<u>.</u>	54
						-			-	_	-		- 7	
		•	•		•	. •		٠	-	_		•	_	

Note. Each - or + indicates the absence or presence of fungal growth from a single test-block.

RESULTS.

From a study of the tabulated results it will be seen that great variation in heat resistance is shown by wood-destroying fungi.

The most resistant of the fungi tested, Lentinus lepideus, was killed in moist wood (1 in. cubes) by treatment for sixty minutes at 65° C.

Comparison of Tables I and II shows that close correlation exists between the times taken to kill a fungus by heat in agar culture and in wood blocks.

STIMMARY

An account is given of experiments made to determine the time, at certain temperatures, required to kill a number of wood-decaying fungi both in agar culture and also in moist wood blocks. The results are given in tabular form.

A new method is outlined for the experimental heat treatment of

wood on a small scale, by the use of rubber bags.

The writer is indebted to Mr W. P. K. Findlay for suggesting this study and for his kind interest in the work, and to Prof. W. Brown for his helpful advice and criticism. He also wishes to tender his thanks to Mr Hatton, Director of the East Malling Research Station, for permission to finish this investigation at the East Malling Laboratory.

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THE PARASITISM OF BOTRYTIS CINEREA PERS. ON AUCUBA JAPONICA THUNB.

By GEORGE TRAPP, M.A., B.Sc., Ph.D. (Botany Department, University of Glasgow)

(With Plates VI and VII)

Introduction

While a number of cuttings of Aucuba japonica Thunb. were being kept in sterile boiling tubes as experimental material for the investigation of a blight of that shrub it was observed that one of the cuttings became affected by a rapid brown rot which, by the end of several days, had killed it completely. As further investigation showed that the disease was due to the parasitic invasion of Botrytis cinerea Pers., it was considered desirable that the symptoms and effects of the disease as well as the life history of the causal agent should be described in detail sufficient to render certain a pathogenicity hitherto unrecorded for this host (1, 2).

ISOLATION OF FUNGUS AND ESTABLISHMENT OF PARASITISM

The specimen on which the attack was first observed (Pl. VI, fig. 1) bore an inflorescence in the cleft between the branches of the forked stem. The fact that other cuttings without flowers after being similarly tubed for several months not only showed no signs of disintegration but had, on the contrary, produced an adventitious root system near the cut end of the stem, which rested on cotton-wool in sterile water at the base of the tube, suggested that there might be a more than fortuitous connection between the presence of an inflorescence and the occurrence of decay. It was, indeed, only at the uncuticularised surface of the nectaries that infection of undamaged twigs could be effected. This is clearly shown (Pl. VI, fig. 1) by the inability of the superficial mycelium to penetrate the intact surface of the, as yet, unattacked portion of the stem below the peduncle. The necrosis had, as indicated (Pl. VI, fig. 1), spread upwards into the younger, more succulent tissues of the limbs of the fork but when a portion of this necrotic tissue was excised after surface sterilisation and transferred on the point of a sterile scalpel to the floral nectaries of another twig the disease first spread downwards into the harder tissues of the older part of the stem (Pl. VI, fig. 2). Pieces of diseased tissues removed with the usual aseptic precautions, and planted on the sterile agar surfaces of poured plates yielded, after twenty-four to forty-eight hours' incubation at 20° C., abundant growth of pure mycelium of characteristic Botrytis form from which subcultures of unquestionable purity were then obtained by hyphal tip transfers. With the fungus from diseased twigs, thus isolated in pure culture, inoculations were carried out on healthy twigs freshly cut and enclosed in sterile boiling tubes. When such a twig was abraded or incised with a sterile instrument following the surface sterilisation of selected areas, and the fungus introduced by implantation on drops of solidified sterile agar placed on these sites, the presence of brown necrotic symptoms was apparent by the end of the first day. Where the mid-rib region of the lamina, the stem apex and the nodes were inoculated (Pl. VI, fig. 3) the signs of fungal invasion, as indicated externally by the appearance of a brown rot, spread out in all directions through vascular and ground tissue alike and had reached the limits shown by the third day (Pl. VI, fig. 3). On re-isolation of the fungus from twigs diseased as a result of artificial inoculation it proved to be morphologically and culturally identical with that originally isolated and subsequently used as inoculum. Control twigs similarly treated, but for the omission of the suspected pathogen, remained perfectly healthy over the protracted period of observation.

It should be mentioned at this point that, in spite of the proved virulence of the parasitism of the fungus for healthy and even vigorously growing tissues of Aucuba, no evidence of disease attributable to this fungus was found in nature nor was it ever isolated from the lesions of Aucuba plants affected by blight or other naturally occurring form of disease. From this it seems that not only is the presence of an exposed or injured surface essential to infection by the fungus but also that the determining factor in its progressive invasion is the concomitance of the moist, enclosed conditions under which the

inoculation experiments were conducted.

PATHOLOGICAL HISTOLOGY

The nature of the invasion was also examined histologically by fixing, in Flemming's strong fluid, portions taken from the externally visible region of demarcation between the brown infected and green healthy tissues and cutting longitudinal sections 4μ thick. Examination of suitably stained sections from the apical part of a stem, including the basal portions of the petioles of the apical leaves, which had been inoculated some distance below the apex, showed that the rot was due to an indiscriminate or parenchymo-vascular attack on the tissues with intracellular penetration of the hyphae. Since the material referred to was in active growth when fungal invasion took place there was little distinction between the rate or mode of progress of the fungus in the vascular tracts and cortex and

pith respectively, except that the hyphae tended to run more longitudinally in the primordial vascular strands, where the septate character of the mycelium could readily be recognised, compared with their more tortuous course in the ground parenchyma. Stained with Flemming's triple stain the tips of the leading hyphae, retaining the safranin more strongly than the older vacuolate segments of the mycelium, showed clearly the intracellular invasive powers of the parasite (Pl. VII, fig. 1). A better stain for showing the general distribution of the pathogen in the host tissues is a solution of 1 per cent. cotton blue in lactophenol. Sections stained for three days in this medium and differentiated for about half that period in the pure solvent (lactophenol) show bright blue fungal cytoplasm and unstained hyphal walls while the walls and cytoplasm of the host cells are almost colourless and the nuclei pale greenish blue (Pl. VII, fig. 2). The central hypha shown invests the less intensely stained circular nucleus at the centre of the cortical host cell; the black spots in the body of the fungus are oil globules stained by the osmic acid of the fixative.

IDENTIFICATION OF THE PATHOGEN

In establishing the identity of the causal agent as Botrytis cinerea Pers. the life history of the organism was briefly worked out in pure culture. Infected tissues from the interior of a diseased plant were sown at opposite sides of a poured plate on the sterile surface of agar prepared with a decoction of Aucuba twigs. Three days later typical hyaline greyish white Botrytis mycelium had grown out from these opposing inocula and covered almost the entire available surface of the agar with spreading hyphae which showed characteristic branchings (Pl. VII, fig. 3). After a further day or so aerial hyphae arose, principally around the margins of the Petri dish and along the junction of opposing growths, and when examined microscopically were seen to consist mainly of stouter brownish conidiophores which bore at their distal ends clusters of typical Botrytis conidia (Pl. VII, fig. 4). On a synthetic medium of sucrose 2 per cent., KNO₃ 0.2 per cent., MgSO₄ 7H₂O and KH₂PO₄ 0.05 per cent. each and washed agar 3 per cent., the mycelium after rapidly covering the available agar surface becomes denser and, probably because of that, appears much whiter. A short time later (five to six days in all) it is observed that at a number of points on the agar surface the hyphae have aggregated to form round, discrete, black sclerotia on the outside of which beadlets of exuded water can be seen (Pl. VII, fig. 5). When transferred to a fresh agar surface these sclerotia immediately germinate to produce an extensive mycelial system which, when it has exhausted the immediately available nutriment or produced a staling depressor, aggregates to form a fresh crop of resistant sclerotia. Sclerotia are also formed at the surface of Aucuba stems killed by the fungus and when mature they become rounded off and fall away. In the dead leaves sclerotia of a different type are found. The fungus here forms irregular patches of pseudo-hypertrophy in the mesophyll the protuberance being towards the abaxial side and its limits often being defined by the course of the larger veins. An even more rapid production of sclerotia could be obtained by inoculating young succulent autoclaved twigs of Aucuba with the sporulating mycelium of Botrytis, when large numbers of sclerotia as well as abundant conidial fructifications were soon evident at the surface of the twigs.*

Healthy twigs of Aucuba japonica Thunb. are actively parasitised by Botrytis cinerea Pers. under conditions of enclosed humidity and provided some susceptible ingress is afforded the pathogen in the form of an unprotected nectary surface or mechanical interruption of the cuticle.

The signs of the disease and its course are described and the nature

of the invasion examined histologically.

In identification, the various phases in the life history of the causal organism have been briefly outlined.

ACKNOWLEDGMENT

The work was carried out during part of the time the author was in receipt of a grant from the Department of Scientific and Industrial Research.

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EXPLANATION OF PLATES VI AND VII PLATE VI

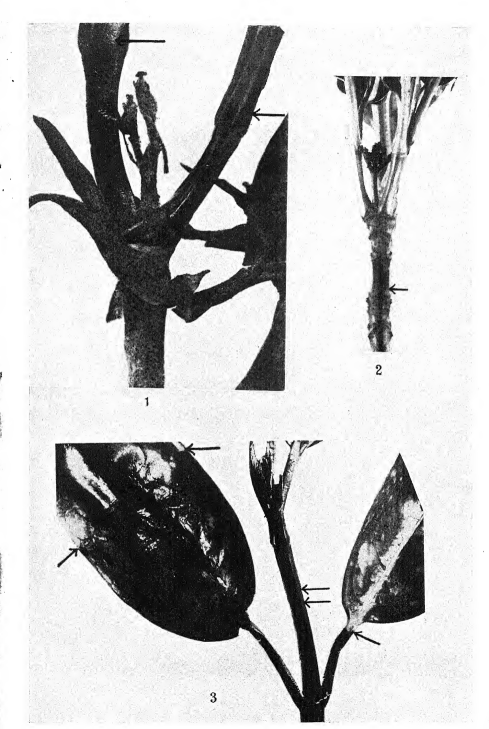
Fig. 1. Experimental cutting of Aucuba from boiling tube. Botrytis infection has taken place through the exposed stigmata and nectaries and the rot has spread from the peduncle into both limbs of the fork as far as the levels indicated. The growing points of the branches have been killed and the necrosis is spreading to the petioles of the apical leaves. x2.

Fig. 2. Infection has occurred as described for fig. 1, but the disease has advanced into the older, more woody part of the stem below the peduncle to the point shown: the younger, more succulent branches, above the junction of inflorescence and main

axis, have not yet been affected. × 3.

Fig. 3. Result, after three days, of artificial inoculations of an Aucuba twig with sporulating mycelium of Botrytis isolated from lesions similar to those depicted in figs. 1 and 2. The artificially induced lesions have spread from the sites of inoculation at (1) the mid-rib of the lower left-hand leaf at the mid-laminal region, (2) the open surface of the petiole of the left-hand apical leaf deprived of its lamina, and (3) the stem just above the lowest node shown. The limits of the general tissue rot are as defined. × 1.

* The sclerotia being produced towards the moister lower end of the twig and the conidia at the drier surface near the top.





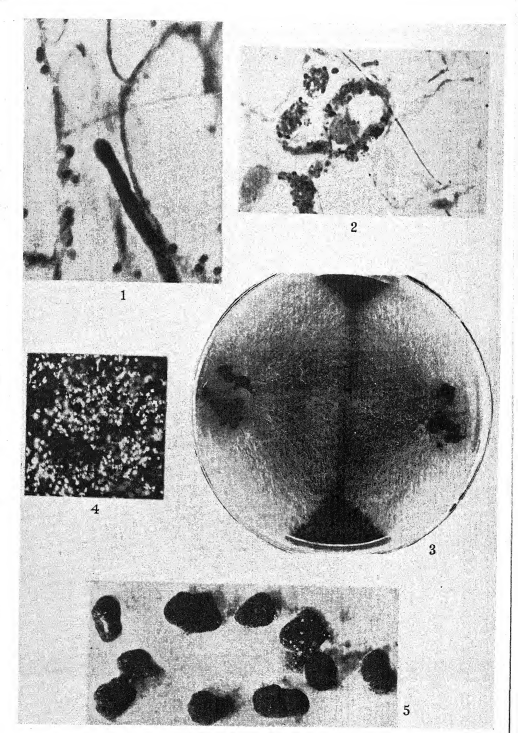




PLATE VII

- Fig. 1. Longitudinal section of an infected Aucuba stem showing the parasitic Botrytis mycelium. The densely granular advance hypha of the invasion is penetrating the host cells of an apical petiole. Stained Flemming's triple stain. ×830.
- Fig. 2. Hypha ramifying in the cells of the killed cortical region of an Aucuba stem (from a section similar to that described for fig. 1). The more lightly stained circular bodies are the host nuclei. Sections from material fixed in Flemming's strong fluid and stained with cotton blue in lactophenol. × 780.
- Fig. 3. Re-isolation of Botrytis in pure culture on agar. Inocula derived from artificially
- infected Aucuba stem lesion (three days old culture). $\times \frac{1}{2}$. Fig. 4. Conidial heads of the aerial mycelium. Note typical clusters of Botrytis conidia. ×и.
- Fig. 5. Group of Botrytis sclerotia standing out from dense white parent mycelium showing surface morphology and droplets of exuded moisture (from six days old subculture on nutrient agar). ×4.
- Figs. 3, 4 and 5 photographed by reflected light.

NOTES ON A TECHNIQUE FOR THE LABORATORY EVALUATION OF PROTECTIVE FUNGICIDES

By R. W. MARSH

(With 1 Text-figure)

THE usefulness of laboratory tests of protective fungicides has recently been pointed out by Montgomery & Moore (1) who have summarised many of the difficulties involved. Montgomery & Moore describe a method of testing based on the distribution of a drop of spray fluid of known volume over a given area on a glass slide. The writer has attempted to carry out parallel tests on slides and on leaves, and for this purpose it has been found necessary to apply the spray by means of an atomiser. The defect of glass slides is that spray fluids may spread too readily on them: this has been remedied by using slides previously covered with cellulose according to the method described by Evans & Martin (2). A brief description follows of the routine adopted in carrying out tests on slides and on leaves, together with a comparison of some of the results obtained.

TESTS ON SLIDES

Glass microscope slides 3 × 1 in. are "cellulosed" by dipping them in a solution of nitro-cellulose in butyl acetate, allowing them to drain and then dry in the laboratory for two or three days. The dry cellulose film is easily detachable in water, so that, in slides that are to be leached, the film must be sealed around the glass by placing small drops of a gum such as Euparal (3) along the edge of the slide.

A standardised method of spraying the slides was adopted using an atomising apparatus of the type described by Evans & Martin (2). The essential feature of the apparatus is the two jets cut from stainless steel tubing of 0.022 in. diameter. The air pressure employed is two atmospheres and the slide is exposed for ten seconds opposite and at a distance of 2 ft. from the spray jet in the plane at right angles to the axis of the spray cone. This exposure gives an even deposit of minute droplets on the sprayed surface at the rate of 0.05 c.c. of spray fluid per square inch.

After being sprayed, the slides are allowed to dry in the laboratory for two days and then, if a leaching treatment is desired, they are submerged for an hour in a litre of fresh rainwater in a glass dish. After leaching, the slides are then again allowed to dry in the

laboratory.

The spores used for a test of the toxicity of the spray deposit are conidia of *Venturia inaequalis*, taken from the youngest visible naturally occurring leaf infections on leaves of Crimson Cox apple. A suspension of these conidia is made in tap water and three drops of the suspension, each of approximately 0.015 c.c., are placed in line at 1 cm. intervals along the middle of the sprayed side of the slide. The concentration of the suspension is such that each drop contains 200–300 conidia. The inoculated slide is then inverted and enclosed over water in a Petri dish. After twenty-four hours in the laboratory a count of spore germination is made; the slide is then returned to the Petri dish and a second count is made after a further twenty-four hours.

TESTS ON LEAVES

Apple leaves of the variety Crimson Cox are employed and in the early part of the season these are obtained by cutting spurs from the tree but from July onwards extension shoots 1 ft. to 1 ft. 6 in. long are taken. A single young leaf about 11 in. long is selected near the tip of the shoot and this leaf is allowed to remain attached to the stem, the others being removed. For the spray treatment, the procedure is exactly as described for the cellulosed slides, the stem bearing the single leaf being supported so that the leaf stands at right angles to the path of the spray with the upper surface of the leaf facing the jet. After the spray application, the stem carrying the treated leaf is placed with the cut end in water and the spray deposit is allowed to dry in the laboratory for forty-eight hours. If the leaf is to be leached it is then subjected to a sprinkling process simulating natural rainfall. In this process, distilled water is allowed to issue at the rate of twelve litres per hour from a fine rose, and the jets thus produced fall approximately a foot on to the leaf upper surface supported in a horizontal position. This treatment is continued for an hour, after which the leaf is again permitted to dry in the laboratory for a further twenty-four hours, the cut end of the stem meanwhile remaining in water.

Before inoculation, the shoot bearing the treated leaf is shortened to about 2 in. and the cut end of the shoot is placed in water in a specimen tube (3×1) in.). Inoculation of the leaf is carried out using the same spore suspension as is employed for the slides, two drops each of 0.015 c.c. being placed on the upper surface, one on either

side of the midrib.

The inoculated leaf is immediately covered by a glass bulb $(2\frac{1}{2})$ in long and 1 in. diameter) having the lower end open. This lower end is then fitted within the top of the collecting tube, water is added to fill the tube and the orifice of the bulb is thus sealed enclosing the

leaf in a moist chamber (see Fig. 1). The whole apparatus is then kept in a humid atmosphere under a bell jar in the laboratory for

twenty-four hours.

fungicidal action.

At the end of this incubation period the small section of the leaf carrying one of the drops is cut out and the leaf then returned to the moist chamber as before. The excised portion is warmed on a slide in a clearing fluid made by mixing equal weights of crystalline chloral hydrate and crystalline phenol, both being first made fluid by heating. In this clearing fluid the leaf fragment becomes translucent in one or two minutes. The preparation is then examined microscopically and a count is made of the total number of spores and of the number germinated. With apple scab

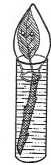


Fig. 1. Apparatus for spore germination tests on leaves. (For description see text.)

conidia the germ tubes are visible without staining.

After a further twenty-four hours the second inoculation drop remaining on the leaf is similarly prepared for a germination count.

It is found that preparations treated with the chloral hydratephenol mixture can be examined without further treatment for approximately a week after clearing.

RESULTS

Table I sets out the results obtained with four types of treatment no spray, lime sulphur, cuprous cyanide and cupric ferrocyanide. The lime sulphur was used at I per cent. strength (by volume), the copper compounds at a concentration equivalent to 2 gm. Cu per litre. The latter materials were supplied by Imperial Chemical Industries, Ltd., in the form of pastes giving excellent dispersion in water. Tests made showed that the dispersing agent used was without

To avoid complexity, the results given in Table I are all taken from surfaces that have not been leached, and the table is planned to facilitate comparison between corresponding treatments on slides and on leaves. While there are certain wide variations in the germination figures relating to the same treatment, the results agree in showing that the spray deposit of cuprous cyanide in these tests was completely fungicidal on slides but of indifferent fungicidal power on leaves. Again, with the spray deposits from cupric ferrocyanide and from lime sulphur, the toxicity shown on leaf surfaces was significantly less than that shown on the cellulosed slides.

In the unsprayed series, the germination on leaves was significantly

Table I. Percentage germinations of apple scab conidia

with le	nation 48 hr.	On leaves		92		53	•	70		53		09						99		5
prayed	Germination after 48 hr.	$^{ m On}$	٠.	89		0		0		0		0						14	•	neans==
On surfaces sprayed with cupric ferrocyanide	Germination after 24 hr.	On leaves		73		28		72				35						09		tween n nt.
On st	Germination after 24 hr.	$\stackrel{'}{ m on}$	٠.	31	•	0		0				0		•				ω		ence be
ıith .	ation 3 hr.	On leaves				33		25				0		47	2	40	24	36		Comparing columns 3 and 4, difference between means=7 Standard error of above difference = 5 i.e. difference is not significant.
On surfaces sprayed with cuprous cyanide	Germination after 48 hr.	On Slides	۰ ،	•		0		0				0		0				0		ns 3 and bove dil lifference
surfaces sprayed v)	On leaves	5 6 .		•	40		65				0		35	S1	43	46	39		column rror of a i.e. d
On su	Germination after 24 hr.	$O_{ m n}$	۰ .			0		0			•	°o		0		•		0		mparing ndard ei
ith	tion hr.	On leaves	ο.		•			90		IO				84	55.	92	46	26		Sta
rayed w phur	Germination after 48 hr.	On slides le						0		47		0		0		•		12	,• ×	= 3.3
On surfaces sprayed with lime sulphur	₹ _	c a s	œ.					29				0		73	78	78	40	44		means
On sur	Germination after 24 hr.	On slides 1	11					0	•			0		0	•			33		between ınt.
çç	ion i	On leaves				00		1(93	98	87	98	74	2	16			84	12.1	nns 1 and 2, difference bet fabove difference i.e. difference is significant.
l surface	Germination after 48 hr.	On Slides le				75	70				81 8		69		89	, .			8.5	nd 2, dil differen rence is
On unsprayed surfaces	₹	On leaves sl	73	84	93	87		83	82			833	70	84	87			82	6.9	mns 1 a of above i.e. diffe
On m	Germination after 24 hr.	On slides le	17	82	70	75	71	8	72		•	ξ,	99	70	27			72	0.6	ing colu d error o
		s 5561	Apr. 26	May		May 16		May 92	Cr (mrii	May 91	tray 3.	Line 14	Trans	Time oo	June 22			Mean	Standard deviation	Comparing columns 1 and 2, difference between means=10 Standard error of above difference = 3. i.e. difference is significant.

better than that on slides in the count taken twenty-four hours after sowing, but no significant difference was shown on the count taken after forty-eight hours. This would suggest a stimulating effect by the leaf sufficient to accelerate germination, but further data are

required to decide this point.

Comparing the sprayed leaves with the sprayed slides, it is possible that the less effective performance of the fungicide on the former is related to a stimulatory effect of the leaf on spore germination but in considering this point the possibility of other influences must be borne in mind. If a reducing substance is produced by the leaf surface then the fungicidal effect of a sulphur-containing spray deposit might, on the hypothesis of Barker (4) be greater on a leaf than on a slide. Conversely, if the fungicidal value of a cuprous salt is dependent on its conversion to the cupric form, a reducing effect on the leaf surface should be prejudicial to the effectiveness of cuprous cyanide. It may be suggested that these biochemical influences of the leaf are outweighed by the factor of the distribution of the spray deposit. The cellulosed slide provides a perfectly plane surface while the leaf presents an epidermis diversified by irregularities, obstructed by hairs, and capable of growth after spray treatment. Therefore while the slide and the leaf may receive precisely the same spray treatment, the spray deposit encountered by the fungus spores in the germination test may be less evenly distributed on the leaf surface than on the slide. Further experiments may show whether this supposition is justified but in the meantime it appears correct to say that in the tests reported above, the living leaf has proved less flattering to the fungicide than has the slide.

SUMMARY

1. A technique is suggested for the laboratory evaluation of pro-

tective fungicides using (a) cellulosed slides, (b) living leaves.

2. Using the same spray treatment on slides and on leaves the spray deposits from cuprous cyanide, from cupric ferrocyanide and from lime sulphur displayed less fungicidal effect on leaves than on slides.

ACKNOWLEDGMENTS

Acknowledgments are made to my colleagues, Messrs A. C. Evans, H. G. H. Kearns and H. Martin, for inspiration and ready assistance on numerous occasions during this investigation.

The cuprous cyanide and cupric ferrocyanide preparations were provided by Imperial Chemical Industries, Ltd., who also contributed to the cost of the spraying apparatus used.

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A NOTE ON THE GROWTH OF THE APPLE SCAB FUNGUS (VENTURIA INAEQUALIS ADERH.) ON BRAMLEY'S SEEDLING APPLES DURING THE WINTER 1934–1935

By W. F. CHEAL, D.I.C.

The breakdown in the resistance of Bramley's Seedling to attacks of the apple scab fungus has become a matter of great economic importance. In certain districts, spraying trials in 1934 showed that two pre-blossom fungicide applications in addition to the usual treatment after the flowering period were necessary in order to obtain a successful scab control: the spray programme for Bramley's equalled the drastic measures required for the most susceptible varieties, e.g. Worcester Pearmain.

Storage problems in 1934 were made very difficult by the large crop, and many tons of Bramley's Seedling apples were kept under outdoor conditions—in clamps, or in orchard boxes stacked and roofed with straw.

It was noticeable late in October of that year that scab colonies on Bramley's apples kept out-of-doors were unusually vigorous; fruit from spraying trials, which had been graded for scab in the previous month had to be placed in a higher category for scab infection. Even in the following month fungal growth appeared to be active, and in December it was decided to make some detailed observations.

Four Bramley's Seedling apples each about 8 oz. in weight, infected with scab, were selected on December 10, 1934, and the positions and dimensions of fifteen scab colonies were recorded. The maximum and minimum diameters of each colony were measured.

The apples were kept in a chip basket placed in a small grass enclosure at Wisbech, and the fungal colonies were measured on December 28, 1934, and again on February 13, 1935.

The figures obtained are shown in Table I.

With three exceptions, growth took place on all the colonies under observation.

The winter 1934-5 was an extremely mild one, and the high temperature for December was outstanding.

The weather conditions permitted scab to develop even on apples stored in the open. The present note therefore confirms the observations of Dr H. Wormald* who found that scab spots may increase in

^{*} J. Min. Agric. September, 1934.

size on Bismarck apples in storage during winter, and this fact emphasises the importance of "pin spot" scab at picking time.

Table I

Apple	Fungus colony	Dec. 10, 1934 mm.	Dec. 28, 1934 mm.	Feb. 13, 1935 mm.
A	i	13×5	14' × 5*5	Fruit bird pecked and rotten
	ii	6×7	7×8	,,
В	i ii	7×6	7×6 6.5×5	7×6
	iii	5×4.5		7×6
	iv	5×5 7×6	5 × 5 8 × 6	5×5 8×7
	v vi	3×4	4 × 5	5×5
~	۸1	4×4	4.5×4	5×5
C	.1	7×9	8 × 10·5	11×9
	.ii	5×4	5.5×5	6×5.5
	iii iv	3×3.5	4.5 × 4 6 × 4.5	5×4·5
		5×4	1.0	6×5.5
	v	7×4	7·75× 4	9×5
D	i	7×7	7 × 7	7×7
	ii	3×3	4 × 4	5×5·5

Average diameter of 13 colonies on apples B, C and D 5.076 mm. (Dec. 10, 1934), 6.23 mm. (Feb. 13, 1935).

REVIEWS

Le Genre Galera (Fr.) Quélet. By ROBERT KÜHNER. (Paul Lechevalier. Paris. Frs. 75.)

The determination of the smaller agarics is beset with difficulties, and monographs are badly needed. The well-known French mycologist Robert Kühner, has given us the results of many years' continuous work on the genus *Galera*. He describes 44 species with many varieties and forms.

The species are grouped into two genera and two sub-genera, including many of the smaller Pholiotas and Naucorias, the microscopic characters of which, in the author's view, place them among the Galeras. We regret the disappearance of Galera as a genus.

The descriptions are full and the microscopic characters are given in great detail. There are no coloured plates, but numerous figures of spores, basidia and cystidia. A key to the species is provided.

The author is to be heartily congratulated on his achievement.

A. A. P.

Flora Agaricina Danica. By JAKOB E. LANGE.

The name of Jakob E. Lange is familiar to all students of the agarics. He has produced a series of brochures that have been published at fairly regular invervals since 1914, dealing chiefly with the microscopic characters of various genera. He has always referred us to his paintings, but as they are in Denmark, few mycologists have had the advantage of consulting them. At last he has made arrangements to have these paintings reproduced under the title Flora Agaricina Danica and the first half of Part I has been published in Copenhagen by the Society for the Advancement of Mycology in Denmark and the Danish Botanical Society. It consists of 16 coloured plates and 40 pages of brief descriptions. Several species are on each plate, and they are splendid examples of the mycologist's art; the colourist and the systematist combining to give a recognisable picture of the species.

The figures of uncommon agarics will be of great assistance to agaricologists, and we hope that the work will receive the recognition it so fully deserves.

A. A. P.

PROCEEDINGS

Meeting held at University College, London, January 19, 1935. *President:* MALCOLM WILSON, D.Sc., F.R.S.E., F.L.S.

L. E. HAWKER. Factors influencing sporulation of *Melanospora* and some other fungi.

A strain of *Melanospora destruens* was found to fruit more freely in the presence of certain other fungi than it did in pure culture. Reduction in the glucose content of the medium or transference from the normal medium to a more dilute one increased to some extent the number of perithecia formed. Media prepared from nutrient liquids in which certain other fungi had been growing were more favourable to perithecial formation than was fresh ("unstaled") medium. Evidence was obtained that the stimulatory effect of certain fungi was due to a combination of three factors, i.e. reduction in food content of the medium, production of "staling" substances by the fungus and production of a definite stimulatory substance or substances. An extract of lentils known to contain accessory substances essential for the growth of *Nematospora Gossypii* stimulated the sporulation of *Melanospora* to a striking extent. The effects of lentil extract were compared with those of certain staled media, under a variety of conditions, and it was suggested that the stimulatory substances concerned were the same or at least very similar.

R. Hull. Investigation of the control of spoilage of processed fruit by Byssochlamys fulva.

The presence or absence of Byssochlamys fulva, a spoilage agent of processed fruit, in any given material was determined by incubating at 30° C. samples of leaves, fruit and straw, collected in plugged sterile tubes filled with hot potato-sucrose agar, acidified to pH3 and heated at 80° for 30 minutes, the ascospores of this fungus being resistant to this temperature.

Positive infections were obtained from diseased and healthy samples collected during several months and from several localities, including Colchester, Kent and Gloucestershire. Elimination of the fungus from the field is therefore evidently

impracticable.

In studying possible control practices in the factory disinfectants were found to be ineffectual, a temperature of 92° C. was necessary to kill the ascospores in $1\frac{1}{2}$ minutes, increase in sucrose up to 20 per cent. made the spores more resistant but at higher concentration the germination rate declined. It is hoped to make further investigations to discover the best way of destroying the fungus within the limits set by the canning process.

J. RAMSBOTTOM. A British species of Kordyana.

An account was given of the genus Kordyana and a new species was described which had been found on Scirpus lacustris in England.

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Meeting held at University College, London, March 16, 1935.

President: MALCOLM WILSON, D.Sc., F.R.S.E., F.L.S.

A. Burges. Mycorrhizal Investigation in Australia.

The widespread nature of fungal infection of a mycorrhizal type was emphasised and a brief account given of the extent to which mycorrhiza had been found in Australian plants. Mention was made of McLennan's work on Lolium and of McLuckie's on the New South Wales Orchids.

Then followed a more detailed account of the mycorrhiza of *Eriostemon Crowei* as described by McLuckie and Burges and of *Lobelia* as described by Fraser.

In *Eriostemon* two types of fungal invasion were recorded, a *Rhizoctonia* infection of the superficial cells and an extensive infection by an arbuscule- and vesicle-forming fungus. The morphological changes seen in the arbuscules and vesicles were described.

In *Lobelia* an unusual mycorrhizal infection is found. Penetration is from rhizomorphic strands and the mycelium develops extensively in the intercellular spaces which are greatly enlarged. The hyphae become filled with oil. Later the hyphae are crushed by the cells and the oil disappears; at the same time oil appears in the cells of *Lobelia*.

D. M. CAYLEY. Spores and spore germination in wild and cultivated mushrooms.

W. P. FINDLAY. Some observations on the fungi occurring in coal mines.

Fungal decay of pitwood is sometimes very rapid in shallow damp mines: this rot occurs mainly in the props used in the permanent roads and in the return airways. The majority of the pit props used in this country are of coniferous timber. The sporophores produced in mines are often abnormal; in many species of Agaricaceae the pileus is greatly reduced and the stipe is elongated and branched, while in the Polyporaceae the sterile tissue is often reduced and the fructification may consist of a pore surface only.

The principal species found attacking softwood props are Poria Vaillantii and other similar species of Poria, Merulius lacrymans, Paxillus panuoides, Lentinus lepideus, Coniophora cerebella, Fomes annosus and Armillaria mellea. On hardwood props Polystictus versicolor and Stereum hirsutum are common.

Preservation of the props by impregnation with a 2 % solution of zinc chloride or of sodium fluoride applied by the "hot and cold" open tank process, greatly increases the life of the props—pitwood treated in this way will last five to ten times as long as the untreated and, therefore, such treatment is definitely an economic proposition.

E. M. WAKEFIELD. An edible species of Volvaria.

Miss Wakefield exhibited a specimen of *Volvaria volvacea*, an uncommon and edible species of a genus until recently regarded as entirely poisonous.

J. RAMSBOTTOM and E. M. WAKEFIELD. Mycological Nomenclature.

Mr J. Ramsbottom read a communication from Mr T. Petch which dealt with the difficulties in applying the International Rules of Botanical Nomenclature in so far as they referred to fungi. It was suggested that much of the trouble would be obviated if Saccardo's Sylloge Fungorum were taken as the starting point.

Mr Ramsbottom then gave a summary of the history of Botanical Nomenclature and indicated the main principles underlying the International Rules. The application of the special articles dealing with fungi was treated in detail and the ambiguities leading to different interpretations pointed out.

Miss E. M. Wakefield followed with a detailed consideration of a number of proposed Nomina Conservanda indicating the points at issue and how that some of the names proposed were those which would be used if the International Rules

were properly applied.

LIST OF MEMBERS

Honorary Members

Bourdot, Abbé H. Saint-Priest-en-Murat per Montmarault, Allier, France. (1935.)

Lister, Miss Gulielma, F.L.S., 871, High Road, Leytonstone, Essex. (1903.) (1924.)

Rea, Mr Carleton, B.C.L., M.A., 6, Barbourne Terrace, Worcester. (1896.) (1918.)

Smith, Miss Annie Lorrain, O.B.E., F.L.S., 44, Stanwick Mansions, Stanwick Road, London, W. 14. (1899.) (1924.)

Ordinary Members

1. Aberdeen, The University Library. (1916.)

2. Adams, Rev. J. H., Landulph Rectory, Hatt, Saltash, Cornwall. (1919.)

3. Adcock, Mr Archie, Upton Road, Norwich. (1921.)

Ainsworth, Mr G. C., B.Sc., Experimental and Research Station, Cheshunt, Herts. (1931.)

5. Ainsworth, Mrs G. C., B.Sc., D.I.C., 3, Cromwell Avenue, Cheshunt, Herts. (1932.)

6. Alberta, University of, Edmonton, Alberta, Canada. (1924.) 7. Alcock, Mrs N. L., F.L.S., M.B.E., Royal Botanic Garden,

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8. Alaily, Mr Y. A. S. El, The Botany School, Cambridge. (1935.)

9. Ashby, Mr S. F., B.Sc., Imperial Mycological Institute, Ferry Lane, Kew, Surrey. (1926.)

10. Barnes, Mr B., D.Sc., Ph.D., F.L.S., Chelsea Polytechnic, London, S.W. 3. (1922.)

11. Barr, Rev. Robert, T.D., M.A., The Manse, Neilston, Renfrewshire. (1918).

12. Barrington, Dr F. J. F., University College Hospital, Medical School, University Street, London, W.C. 1. (1901.)

13. Bartlett, Mr A. W., M.A., M.Sc., F.L.S., Department of Botany, Armstrong College, Newcastle-on-Tyne. (1920.)

14. Bates, Mr G. R., c/o British South Africa Company, Mazoe Citrus Estate, Mazoe, S. Rhodesia. (1930.)

15. Bates, Mrs L. F., B.Sc., 36, Gainsborough Gardens, Golders Green, London, N.W. 11. (1921.)

16. Beardslee, Mr H. C., Perry, Ohio, U.S.A. (1933.)

17. Beaumont, Mr Albert, M.A., Seale-Hayne Agricultural College, Newton Abbot, Devon. (1924.)

18. Bellchambers, Mr A., The Gardens, Eaton Hall, Chester. (1932.)

19. Bewley, Mr W. F., D.Sc., Experimental and Research Station, Cheshunt, Herts. (1922.)

20. Biffen, Professor Sir Rowland H., M.A., F.R.S., 136, Huntingdon Road, Cambridge. (1899.)

Biologist, Plant Research Laboratory, Horticultural Gardens,

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22. Birmingham Natural History and Philosophical Society, c/o G. T. Calvert, Esq., Hon. Librarian, Avebury House, 55, Newhall Street, Birmingham. (1920).

23. Bisby, Mr Guy R., Ph.D., Manitoba Agricultural College,

Winnipeg, Canada. (1921.)

24. Blackman, Professor V. H., M.A., F.R.S., Imperial College of Science, South Kensington, London, S.W. 7. (1900.)

25. Blackwell, Miss E. M., M.Sc., Botanical Department, Royal Holloway College, Englefield Green, Surrey. (1917.)

26. Bolas, Mr B. D., M.Sc., 20, Cambridge Gardens, Winchmore Hill, London, N. 21. (1924.)

27. Bonn, Germany, Institut für Pflanzenkrankheiten, Nuss-Allee 9. (1931.)

28. Borthwick, Professor A. W., O.B.E., D.Sc., Forestry Department, The University, Aberdeen. (1911.)

29. Boston, The Mycological Club, Horticultural Hall, Boston, Mass., U.S.A. (1926.)

30. Bradshaw, Mr F., M.A., D.Sc., Armstrong College, Newcastle-

on-Tyne. (1928.)

31. Braid, Professor K. W., B.A., B.Sc., West of Scotland Agricultural College, 6, Blythswood Square, Glasgow. (1922.)

32. Brazier, Mr E., Ty'n-y-gongl, Love Lane, Stourbridge. (1921.) 33. Brenchley, Mr G. H., B.A., Clare College, Cambridge. (1925.)

34. Brett, Miss M., M.Sc., Ph.D., Northern Polytechnic, Holloway

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35. Brierley, Professor W. B., D.Sc., F.R.A.I., F.L.S., Department of Agricultural Botany, The University, Reading. (1919.)

36. Brinton, Mr R. E. B., 68, Woodstock Avenue, Golders Green,

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37. Brisbane, The Director, Bureau of Sugar Experiment Stations, Department of Agriculture and Stock, Queensland, Australia. (1930.)

38. British Museum, The Trustees of, Cromwell Road, South

Kensington, London, S.W. 7. (1914.)

39. Brooks, Professor F. T., M.A., F.R.S., F.L.S., The Botany School, Cambridge. (1907.)

40. Brown University, Library, East Side Station, Providence, R.I., U.S.A. (1920.)

41. Brown, Professor W., M.A., D.Sc., Imperial College of Science, South Kensington, London, S.W. 7. (1922.)

42. Bruxelles, Jardin Botanique de l'État, c/o M. P. van Aerdschot. (1911.)

43. Buckley, Mr W. D., "Lynmouth", 2 Curzon Street, Slough. (1916.)

44. Buddin, Mr Walter, M.A., Laboratory of Plant Pathology, University of Reading, 7, Redlands Road, Reading. (1921.)

45. Buller, Professor A. H. R., D.Sc., Ph.D., F.R.S., University of Manitoba, Winnipeg, Canada. (1911.)

46. Bunting, Mr R. H., F.L.S., 3 Stanton Court, Weymouth. (1921). 47. Burges, Mr N. A., The Botany School, Cambridge. (1935.)

48. Burr, Mr S., M.Sc., Department of Agriculture, The University, Leeds. (1924.)

49. Butler, Mr E. J., C.I.E., C.M.G., D.Sc., M.B., F.R.S., F.L.S., Agricultural Research Council, 6 A, Dean's Yard, London, S.W. 1. (1920.)

50. Caldwell, Mr J., B.Sc., Ph.D., Department of Botany, University College, Exeter. (1932.)

51. Cambridge, The Botany School. (1920.)

52. Campbell, Mr A. H., B.Sc., Ph.D., Department of Botany, The University, Bristol. (1934.)

53. Carne, Mr W. M., F.L.S., Australia House, Strand, London, W.C. 2. (1928.)

54. Carr, Professor J. W., M.A., F.L.S., Mapperley Edge, Private Road, Sherwood, Nottingham. (1896.)

55. Carrothers, Mr. E. N., 7, Fitzwilliam Street, Belfast, N. Ireland, (1925.)

56. Carter, Miss F. M., Ph.D., Botanical Department, The University, Edgbaston, Birmingham, 15. (1934.)

57. Cartwright, Mr K. St G., M.A., F.L.S., The Red House, Kingston Blount, Oxford. (1913.)

58. Castellani, Sir Aldo, M.D., 23 Harley Street, London, W. 1. (1922.)

59. Cayley, Miss Dorothy M., John Innes Horticultural Institution, Mostyn Road, Merton Park, London, S.W. 19. (1913.)

60. Charles, Miss Vera K., United States Department of Agriculture, Bureau of Plant Industry, Washington D.C., U.S.A. (1933.)

61. Chaudhuri, Mr H., M.Sc., Ph.D., University of the Punjab, Lahore, India. (1920.)

62. Cheal, Mr W. F., Savile House, Queen's Road, Wisbech, Cambs. (1927.)

63. Chesters, Mr C. G. C., Botanical Department, The University, Edgbaston, Birmingham. (1930.)

64. Ciferri, Professor Dr R., Assistant Director, Laboratorio Crittogamico, Casella Postale 165, Pavia, Italy. (1926.)

65. Clapham, Mr A. R., M.A., Ph.D., Department of Botany, The University, Oxford. (1931.)

66. Cleland, Mr J. Burton, M.D., Professor of Pathology, Univer-

sity of Adelaide, South Australia. (1918.)

67. Clouston, Mr D., M.A., B.Sc. (Agr.), North of Scotland College of Agriculture, Crown Mansions, 41, Union Street (2nd Floor), Aberdeen. (1931.)

68. Colston, Miss B., B.Sc., Ph.D., The University, Manchester.

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69. Connecticut Agricultural Experiment Station, New Haven, Connecticut, U.S.A. (1929).

70. Cook, Mr W. R. I., B.Sc., Ph.D., Department of Botany, University College, Newport Road, Cardiff. (1924.)

71. Cooke, Mr G. J., 143, Newmarket Road, Norwich. (1933.)

72. Cooper, Miss Charlotte A., California Lane, Bushey Heath, Herts. (1911.)

73. Cooper, Mrs V. Astley, The Tors, Knowle, Fareham, Hants.

(1921.)

74. Cornell University, The Library, New York State College of Agriculture, Ithaca, N.Y., U.S.A. (1920.)

Corner, Mr E. J. H., M.A., F.L.S., Assistant Director, Botanic Gardens, Singapore, Straits Settlements. (1924.)

76. Cotton, Mr Arthur D., O.B.E., F.L.S., Keeper, Herbarium, Royal Botanic Gardens, Kew, Surrey. (1902.)

77. Cunningham, Mr G. H., Ph.D., Plant Research Station, Box 442, Palmerston North, New Zealand. (1922.)

78. Curtis, Miss Kathleen M., M.A., D.Sc., D.I.C., F.L.S., Mycologist, Biological Department, Cawthron Institute of Scientific Research, Nelson, New Zealand. (1917.)

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82. Davies, Mr D. W., B.Sc., Adviser in Mycology, Agricultural Buildings, University College of Wales, Aberystwyth. (1923).

83. Davis, Mr J. Jefferson, B.S., M.D., University of Wisconsin, Madison, Wis., U.S.A. (1921.)

84. Day, Mr W. R., B.A., B.Sc., Imperial Forestry Institute, Oxford. (1928.)

85. Deacon, Dr G. E., Brundall, Norwich. (1933.)

86. Dehra Dun, The Forest Botanist, Forest Research Institute and College, U.P., India. (1929.)

87. Deighton, Mr F. C., M.A., Mycologist, Department of Lands and Forests, Freetown, Sierra Leone, West Africa. (1925.) 88. Dennis, Mr R. W. G., Ph.D., West of Scotland Agricultural

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89. Dickinson, Mr S., School of Agriculture, Cambridge. (1921.) 90. Dobbs, Mr C. G., B.Sc., Botanical Department, King's College, Strand, London, W.C. 2. (1933.)

91. Dobinson, Mr H., 166, Piccadilly, London, W. 1. (1932.)

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93. Dowson, Mr W. J., M.A., D.Sc., The Botany School, Cambridge. (1920.)

94. Duncan, Mr J. T., London School of Hygiene and Tropical Medicine, Keppel Street, London, W.C. 1. (1930.)

95. Edwards, Mr W. H., Belle Vue, Barline, Beer, Devon. (1896.)

96. Elliott, Mr W. T., D.D.S., L.D.S., F.L.S., F.Z.S., Arden Grange, Tanworth-in-Arden, Warwickshire. (1913.)

97. Elliott, Mrs J. S. Bayliss, D.Sc. (B'ham), B.Sc. (Lond.), Arden Grange, Tanworth-in-Arden, Warwickshire. (1911.)

98. Ellis, Mr E. H., B.Sc., Gramarye, Farley Green, Guildford, Surrey. (1936.)

99. Ellis, Miss E. M., St Hugh's College, Oxford. (1930.)

100. Ellis, Mr Holmes, F.R.M.S., 108, Birtwistle Avenue, Colne, Lancs. (1927.)

101. Emoto, Dr Y., Biological Department, Peers' College (Gakushuin), Mejiromachi, Tokyo, Japan. (1929.)

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103. Exeter, Librarian, University College of the South-West of England. (1926.)

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106. Findlay, Mr W. P., B.Sc., A.R.C.S., Courte Falaise, Sevenoaks,

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107. Finlayson, Mr Raymond A., F.L.S., Official Seed Testing Station, Huntingdon Road, Cambridge. (1910.)

108. Fisher, Mr S. D. P., Sackville Street, Leeds. (1930.)

109. Fitzpatrick, Professor H. M., Ph.D., 220, Bryant Avenue, Ithaca, New York, U.S.A. (1935.)

110. Forwood, Mr R., Minett, Muskoka, Ontario, Canada.

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111. Fountain, Mr A. S., F.R.M.S., 55, Moorgate, Rotherham, Yorks. (1934.)

112. Gadd, Mr C. H., D.Sc., Tea Research Institute, Nuwara Eliya, Ceylon. (1921.)

113. Galloway, Mr L.D., Imperial Institute of Agricultural Research, Delhi, India. (1928.)

114. Gardner, Capt. Frederic, c/o Barclays Bank, Jersey, C.I. (1898.)

115. Garside, Mr S., M.Sc., F.L.S., Botanical Department, Bedford College, Regent's Park, London, N.W. I. (1922.)

116. Gates, Professor R. R., D.Sc., Ph.D., F.R.S., F.L.S., King's College, Strand, London, W.C. 2. (1921.)

117. Ghamrawy, Mr Ali K., 39 Monirah Street, Cairo, Egypt. (1932.)

118. Gibson, Miss C. J., B.A., 27, Banbury Road, Oxford. (1933.)

119. Gilbert, Dr E. M., Botanical Department, University of Wisconsin, Madison, Wis., U.S.A. (1922.)

120. Gilbert, M. E., Docteur en Pharmacie, 6, Rue de Laos, Paris (15^e), France. (1924.)

121. Gisborne, Mr J. H., Keble College, Oxford. (1932.)

122. Glasstone, Mrs V. F. C., B.A. (Oxon.), 15, Northumberland Road, Sheffield. (1930.)

123. Glynne, Miss Mary D., M.Sc., F.L.S., Rothamsted Experi-

mental Station, Harpenden, Herts. (1932.)

124. Gorman, Mr M. J., A.R.C.Sc.I., Albert Agricultural College. Glasnevin, Dublin. (1925.)

125. Gould, Mr F. G., Woodrising, Trapps Hill, Loughton, Essex. (1918.)

126. Green, Col. C. Theodore, A.M.S., M.R.C.S., L.R.C.P., F.L.S., 31, Shrewsbury Road, Birkenhead. (1901.)

127. Green, Miss E., M.Sc., 15, Gower Street, London, W.C. 1.

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129. Gregor, Mrs M. J. F., Ph.D., Royal Botanic Garden, Edinburgh. (1927.)

130. Gregory, Mr P. H., Ph.D., Seale Hayne Agricultural College, Newton Abbot, Devon. (1930.)

131. Grieve, Mr B. J., M.Sc., Botanical Department, Melbourne University, Carlton, Victoria, Australia. (1931.)

132. Grinling, Mr C. H., B.A., 71, Rectory Place, Woolwich,

London, S.E. 18. (1913.)

133. Grove, Miss Jessica H., 14, The Tything, Worcester. (1927.)

134. Gwynne-Vaughan, Professor Dame Helen, G.B.E., D.Sc., LL.D., F.L.S., 93, Bedford Court Mansions, London, W.C. I. (1906.)

135. Hanna, Mr W. F., M.Sc., Dominion Rust Research Laboratory,

Agricultural College, Winnipeg, Canada. (1925.)

136. Hansford, Mr C. G., M.A., F.L.S., Mycologist, Department of Agriculture, Kampala, Uganda. (1921.)

137. Harley, Mr J. L., Shrublands, Hethersett, Norfolk. (1932.) 138. Harris, Mr G. C. M., 148, Divinity Road, Oxford. (1934.)

139. Harris, Mr R. V., B.Sc., A.R.C.S. Horticultural Research Station, East Malling, Kent. (1924.)

140. Harrison, Mr T. H., D.Sc. Hawkesbury Agricultural College, Richmond, N.S. Wales, Australia. (1931.)

141. Harvard University, The Library, Cambridge, Mass., U.S.A. (1923.)

142. Hastings, Mr Somerville, M.S., F.R.C.S., 43, Devonshire Street, Portland Place, London, W. 1. (1913.)

143. Hawker, Miss L. E., Botanical Department, Imperial College of Science, London, S.W. 7. (1934.)

144. Heim, M. Roger, Sous-Directeur au Muséum d'Histoire Naturelle, 11, Rue de Médicis, Paris (6e), France. (1930).

145. Heimbeck, Mrs Louise, Brosoe, Levanger, Norway. (1923.)

146. Hemmi, Dr Takewo, Phytopathological Institute, Department of Agriculture, Kyoto Imperial University, Kyoto, Japan. (1923.)

147. Hereford, Mr E. H., 131, Queen Victoria Street, London, E.C. 4.

148. Hickman, Mr C. J., Research Station, Long Ashton, nr. Bristol. (1935.)

149. Hildyard, Mr F. W., 1, Lichfield Road, Kew, Surrey. (1913.)

150. Holden, Professor H. S., D.Sc., F.L.S., Botanical Department, University College, Nottingham. (1923.)

151. Honolulu, Association of Hawaiian Pineapple Canners, P.O. Box 3166, Hawaii. (1929.)

152. Honolulu, The Library, Experimental Station, S.P.A., Box 411, Hawaii. (1920.)

153. Hopkins, Mr J. C. F., D.Sc., A.I.C.T.A., P.B. 74B, Salisbury, S. Rhodesia. (1930.)

154. Horne, Mr A. S., D.Šc., F.L.S., F.G.S., Botanical Department, Imperial College of Science, South Kensington, London, S.W. 7. (1921.)

155. Howard, Mr H. J., F.R.M.S., F.L.S., "Lingfield", 6, College Road, Norwich. (1918.)

156. Hubbard, Miss M. D., B.Sc., Department of Botany, University College of Wales, Aberystwyth. (1933.)

157. Hughes, Mr G. C., Priory Road, Bicester. (1898.)

158. Hughes, Mr J. S., M.A., University Observatory, Oxford. (1927.)

159. Hull, The Librarian, Botanical Department, University College. (1929.)

160. Humphrey, Dr C. J., United States Department of Agriculture, Soil Conservation Service, Safford, Arizona, U.S.A. (1921.)

161. Hurrell, Mr H. E., 60, Albany Road, Great Yarmouth. (1921.)

162. Hurst, Mr C. P., Landulph Rectory, Saltash, Cornwall. (1928.)

163. Ingold, Mr C. T., M.Sc., Ph.D., Department of Botany, The University, Reading. (1935.)

164. Iowa, The Library, State University of Iowa, Library Annex, Iowa City, U.S.A. (1923.)

165. Iowa State College, Library, Ames, Iowa, U.S.A. (1927.)

166. Issatchenko, Professor B. L., Directeur du Jardin Botanique, Leningrad, Russia. (1923.)

167. John Crerar Library, 86, East Randolph Street, Chicago, Illinois, U.S.A. (1929.)

168. John Innes Horticultural Institution, Mostyn Road, Merton Park, London, S.W. 19. (1924.)

169. Johnson, Mr J. W. Haigh, M.Sc., F.I.C., F.L.S., Walton, nr Wakefield. (1919.)

170. Jones, Mr G. H., M.A., Plant Protection Section, Ministry of Agriculture, Cairo, Egypt. (1922.)

171. Jørstad, Mr Ivar, Statsmykolog, Botanisk Museum, Oslo, Norway. (1923.)

172. Keay, Miss M. A., M.A. (Cape Town), The Botany School, Cambridge. (1935.)

173. Keissler, Dr Karl, Direktor d. Botanischen Abteilung, Naturhistorisches Museum, Burgring 7, Wien 1/1, Austria. (1924.)

174. Kelly, Dr Howard A., 1418, Eutaw Place, Baltimore, Md., U.S.A. (1921.)

175. King, Miss M. E., B.A., The Botany School, Cambridge. (1935.)

176. Klika, Mr Bohumil, Hálkova 37, Prague, Vrsovice 553, Czechoslovakia. (1926.)

177. Knight, Mr H. H., M.A., The Lodge, All Saints' Villas,

Cheltenham. (1914.) 178. Kuala Lumpur, F.M.S., The Director of Agriculture, Straits Settlements, and Federated Malay States. (1930.)

179. Lamb, Mr I. M., B.Sc., 18, Duke's Avenue, Kingston-on-Thames, Surrey. (1934.)

324

- 180. Lampitt, Mr L. H., D.Sc., F.I.C., Thornlea, Mount Park, Harrow, Middlesex. (1925.)
- 181. Leach, Mr R., B.A., Agricultural Department, Mlanje, Nyasaland. (1929.)
- 182. Leicester, The Museum, City of Leicester. (1923.)
- 183. Linder, Dr D., Farlow Herbarium, Harvard University, 20, Divinity Avenue, Cambridge, Mass., U.S.A. (1935.)
- 184. Line, Mr James, M.A., School of Agriculture, Cambridge. (1921.)
- 185. Linnean Society, The, Burlington House, Piccadilly, London, W. 1. (1919.)
- 186. Lloyd Library, The, 309, West Court Street, Cincinnati, Ohio, U.S.A. (1907.)
- 187. Loader, Miss F. M., B.Sc., Botanical Department, University College, Southampton. (1927.)
- 188. Lowndes, Mr A. G., M.A., F.L.S., Marlborough College, Marlborough, Wilts. (1922.)
- 189. Lütjeharms, Mr W. J., Assistent aan's Rijks Herbarium, Leiden, Holland. (1930.)
- 190. McDonald, Mr J., D.F.C., B.Sc., F.L.S., Senior Plant Pathologist, P.O. Box 338, Nairobi, Kenya Colony, East Africa. (1923.)
- 191. McLennan, Dr Ethel I., Botanical Department, Melbourne University, Carlton, Victoria, Australia. (1926.)
- 192. Madras University Library, Senate House, Triplicane, Madras, South India. (1925.)
- 193. Maire, M. René, D.Šc., F.M.L.S., Professeur à la Faculté des Sciences de l'Université, Algiers, Algeria, N. Africa. (1907.)
- 194. Marsh, Mr R. W., M.A., Research Station, Long Ashton, Bristol. (1923.)
- 195. Masefield, Mr G. B., c/o Department of Agriculture, Entebbe, Uganda. (1932.)
- 196. Mason, Mr E. W., M.A., M.Sc., F.L.S., Imperial Bureau of Mycology, 17, Kew Green, Kew, Surrey. (1921.)
- 197. Mason, Mrs E. W., Inglenook, 63, King's Road, Richmond, Surrey. (1922.)
- 198. Mason, Mr F. A., F.R.M.S., M.S.P.A., 29, Frankland Terrace, Leeds. (1912.)
- 199. Matthews, Professor J. R., M.A., F.L.S., Department of Botany, The University, Old Aberdeen. (1921.)
- 200. Mehta, Professor K. C., Ph.D., Department of Biology, Agra College, Agra, U.P., India. (1921.)
- 201. Melville, Mr R., B.Sc., Ph.D., 5, Courtway, Twickenham, Middlesex. (1933.)

202. Metcalfe, Mr C. R., B.A., Ph.D., Jodrell Laboratory, Royal Botanic Gardens, Kew, Surrey. (1926.)

203. Michigan Agricultural College Library, East Lansing, Michigan, U.S.A. (1924.)

204. Miller, Professor J. H., B.S., M.S., Ph.D., University of Georgia, Athens, Ga., U.S.A. (1930.)

205. Missouri, The Botanical Garden, St Louis, Mo., U.S.A. (1902.)

206. Mitra, Mr M., M.Sc., Ph.D., D.I.C., Assistant Mycologist, Imperial Institute of Agricultural Research, Delhi, India. (1928.)

207. Miyabe, Dr Kingo, Professor Emeritus of Botany, Hokkaido Imperial University, Sapporo, Japan. (1919.)

208. Montague, Mrs A., Penton, Crediton, N. Devon. (1898.)

209. Montreal, Canada, Faculté des Sciences, Institut Botanique, Université de Montréal. (1932.)

210. Moore, Mr W. C., M.A., Ministry of Agriculture, Pathological Laboratory, Milton Road, Harpenden, Herts. (1922.)

211. Morgan, Dr G., Ashley-Hatton, Dyke Road Avenue, Brighton. (1928.)

212. Morris, Mr L. E., c/o Eton College, Windsor, Berks. (1924.)

213. Muller, Dr H. R. A., Institut voor Plantenziekten, Buitenzorg, Java. (1932.)

214. Murphy, Professor P. A., Sc.D., A.R.C.Sc.I., M.R.I.A., Department of Plant Pathology, Albert Agricultural College, Glasnevin, Dublin, N.W. 3. (1924.)

215. Murray, Mr G. H., F.E.S., Director of Agriculture, Rabaul, New Britain, Territory of New Guinea, via Australia. (1921.)

216. Muskett, Mr A. E., M.Sc., A.R.C.S., Queen's University, Belfast, Northern Ireland. (1923.)

217. Nannfeldt, Dr J. A., Sturegatan 11, Uppsala, Sweden. (1932.)

218. National Collection of Type Cultures, Curator, Lister Institute, Chelsea Gardens, London, S.W. 1. (1921.)

219. National Museum of Wales, Cardiff. (1924.)

220. Nattrass, Mr R. M., B.Sc. (Ágric.), Ph.D., Department of Agriculture, Nicosia, Cyprus. (1925.)

221. Nebraska, The Library, University of, Lincoln, Nebr., U.S.A. (1924.)

222. Nederlandsche Mycologische Vereeniging, c/o Mr A. C. S. Schweers, Nassaulaan 17, Alkmaar, Holland. (1920.)

223. Nelson, Miss M. G., M.A., Botanical Department, The University, Oxford. (1932.)

224. New York Botanical Garden, Bronx Park, New York, U.S.A. (1904.)

225. Nicholson, Mr W. E., F.L.S., 50, St Anne's Crescent, Lewes. (1913.)

- 226. Noel, Miss E. F., F.L.S., 37, Burnham Court, Queen's Road, London, W. 2. (1913.)
- 227. North Carolina, Library, University of, Chapel Hill, North Carolina, U.S.A. (1920.)
- 228. Nursery and Market Garden Industries' Development Society, Ltd., Experimental and Research Station, Cheshunt, Herts. (1922.)
- 229. O'Connor, Mr P., Ph.D., B.Sc., A.R.C.Sc.I., National Museum, Dublin. (1925.)
- 230. Ogilvie, Mr L., M.A., M.Sc., Research Station, Long Ashton, nr Bristol. (1922.)
- 231. Oke, Mr Alfred William, B.A., F.G.S., F.L.S., 32, Denmark Road, Hove, Sussex. (1908.)
- 232. Ontario Agricultural College, Library, Guelph, Ontario, Canada. (1920.)
- 233. Oort, Dr A. J. P., Ericalaan 5, Wageningen, Holland. (1935.)
- 234. Osborn, Professor T. G. B., D.Sc., F.L.S., Botanical Department, The University, Sydney, N.S.W., Australia. (1910.)
- 235. Ottawa, Óntario, Canada, The Library, Geological Survey. (1926.)
- 236. Padwick, Dr G. Watts, Jealott's Hill Agricultural Research Station, Bracknell, Berks. (1936.)
- 237. Page, Miss W. M., M.Sc., 19, Ledam Buildings, Bourne Estate, Holborn, London, E.C. 1. (1921.)
- 238. Park, Mr M., Department of Agriculture, Peradeniya, Ceylon. (1929.)
- 239. Parke Ďavís & Co., Medical Research Library, P.O. Box 488, Detroit, Michigan, U.S.A. (1920.)
- 240. Parker, Professor C. S., Department of Botany, Howard University, Washington, D.C., U.S.A. (1932.)
- 241. Pearson, Mr Arthur A., F.L.S., 59, Southwark Street, London, S.E. 1. (1911.)
- 242. Peklo, Dr Jaroslav, Professor of Applied Botany, Bohemian Technical University, Charles Square, Prague II, Czechoslovakia. (1924.)
- 243. Perthshire Society of Natural Science, c/o Mr James Winter (Hon. Treasurer), 35, George Street, Perth. (1919.)
- 244. Petch, Mr T., B.A., B.Sc., North Wootton, King's Lynn, Norfolk. (1911.)
- 245. Pethybridge, Mr G. H., Ph.D., B.Sc., F.L.S., Ministry of Agriculture, Pathological Laboratory, Milton Road, Harpenden, Herts. (1919.)
- 246. Peyronel, Dr Benjamino, R. Istituto Sup. Agrario e Forestale, Piazzale del Re, Firenze, Italy. (1932.)

247. Philadelphia, The Academy of Natural Sciences of Philadelphia, Nineteenth and The Parkway, Phil., U.S.A. (1925.)

248. Phillips, Dr H. H., 6, St John's Road, Penge, London, S.E. 10. (1923.)

249. Ping, Mr A. Wentworth, M.A., "St Olave's", Clifton, York. (1926.)

250. Potter, Rev. M. C., Sc.D., M.A., F.L.S., Corley Croft, New Milton, Hants. (1896.)

251. Preston, Mr N. C., B.Sc., Harper Adams Agricultural College, Newport, Salop. (1920.)

252. Pretoria, South Africa, The Chief, Division of Plant Industry (91403), Department of Agriculture. (1922.)

253. Purdue University, Library, Agricultural Experiment Station, Lafayette, Ind., U.S.A. (1931.)

254. Pusa, Imperial Mycologist, Imperial Agricultural Research Institute, Delhi, India. (1921.)

255. Ramsbottom, Mr J., O.B.E., M.A., F.L.S., British Museum (Nat. Hist.), Cromwell Road, South Kensington, London, S.W. 7. (1910.)

256. Ray, Miss Anne, Penarwyn, Gorran Haven, Gorran, Cornwall. (1929.)

257. Rayner, Dr M. Cheveley (Mrs Neilson Jones), Bedford College for Women, Regent's Park, London, N.W. 1. (1921.)

258. Rea, Miss M. W., M.Sc., Salem House, Sydenham, Belfast, Northern Ireland. (1920.)

259. Rees, Mr John, M.Sc., Adviser in Agricultural Botany, University College, Cardiff. (1929.)

260. Reichert, Dr Israel, Jewish Agency for Palestine, Agricultural Experiment Station, P.O.B. 15, Rehoboth, Palestine. (1924.)

261. Rhind, Mr Donald, B.Sc., Economic Botanist, Department of Agriculture, Agricultural College, Mandalay, Burma. (1922.)

262. Rhodes, Miss Mabel, Lister Institute, Chelsea Gardens, London, S.W. 1. (1921.)

263. Robson, Mr R., M.Sc., F.Z.S., East Hanningfield, Chelmsford, Essex. (1914.)

264. Rothamsted Experimental Station, Department of Mycology, Harpenden, Herts. (1923.)

265. Rutgers College and State University of New Jersey, Library, New Brunswick, New Jersey, U.S.A. (1922.)

266. St Paul, Minnesota, U.S.A., The Library, Department of Agriculture, University Farm. (1920.)

267. Salisbury, Professor E. J., D.Sc., F.R.S., F.L.S., Botanical Department, University College, Gower Street, London, W.C. 1. (1921).

268. Salmon, Professor E. S., F.L.S., South-Eastern Agricultural College, Wye, Kent. (1922.)

269. Sampson, Miss K., M.Sc., University College, Aberystwyth, North Wales. (1920.)

270. Samuel, Mr Geoffrey, M.Sc., Department of Plant Pathology, Rothamsted Experimental Station, Harpenden, Herts. (1923.)

271. Scott, Mr W. W., 13, Bishop's Road, Highgate, London, N. 6. (1922.)

272. Searle, Mr G. Odell, B.Sc. (Agric.), Flax Research Institute, Flitcham Abbey, nr. King's Lynn, Norfolk. (1920.)

273. Seth, Mr N. L., B.Sc., Ph.D., D.I.C., Agricultural College, Mandalay, Burma. (1930.)

274. Sharples, Mr A., A.R.C.S., D.I.C., c/o Messrs Grindlay & Co., Parliament Street, London, S.W. 1. (1924.)

275. Shaw, Mr F. J. F., D.Sc., A.R.C.S., F.L.S., Imperial Agricultural Research Institute, Delhi, India. (1920.)

276. Shear, Dr C. L., U.S. Department of Agriculture, Bureau of Plant Industry, Washington, D.C., U.S.A. (1930.)

277. Small, Mr W., M.B.E., M.A., Ph.D., B.Sc., Director, Department of Agriculture, Zomba, Nyasaland. (1915.)

278. Smith, Mr Alexander, M.A., Ph.D., Ministry of Agriculture, Pathological Laboratory, Milton Road, Harpenden, Herts. (1924.)

279. Smith, Miss K. E., The Quarry, Lutterworth Road, Nuneaton. (1913.)

280. Smith, Professor Noel J. G., Ph.D., B.Sc., Botany Department, Rhodes University College, Grahamstown, S. Africa. (1924.)

281. Smith, Mr Rupert, 38, Greenhill Gardens, Edinburgh. (1927.)

282. South London Botanical Institute, 323, Norwood Road, Tulse

Hill, London, S.E. 24. (1921.)
283. Stakman, Professor E. C., University of Minnesota, Department of Agriculture, University Farm, St Paul, Minn., U.S.A. (1922.)

284. Statham, Miss E. M., 2, Westbrook Road, Blackheath, London, S.E. 3. (1926.)

285. Stationery Office, H.M., Superintendent of Publications, Book Dept., Westminster, S.W. 1. (4 subscriptions.) (1920.)

286. Stephens, Miss E. L., B.A., Department of Botany, University of Cape Town, Cape Town, South Africa. (1928.)

287. Stephens, Miss F. L., M.Sc., Department of Botany, British Museum (Natural History), Cromwell Road, South Kensington, London, S.W. 7. (1930.)

288. Steyaert, M. R. L., Station de Sélection Cotonnière de Bambesa, District des Uélés, Belgian Congo. (1931.)

289. Stirrup, Mr H. H., M.Sc., Midland Agricultural College,

Sutton Bonington, Loughborough. (1922.)

290. Storey, Mr H. H., M.A., Ph.D., East African Agricultural Research Institute, Amani, Tanganyika Territory, East Africa. (1922.)

291. Sutherland, Mr G. K., M.A., D.Sc., F.L.S., "Bremhill", 21,

Combe Park, Bath, Somerset. (1914.)

202. Swanton, Mr E. W., M.B.E., A.L.S., Educational Museum, Haslemere, Surrey. (1899.)

293. Swedish Academy of Sciences, Royal, Stockholm, Sweden. (1919.)

294. Sydney, Australia, The Librarian, University of. (1922.)

295. Sydow, Herr H., Luitpoldstrasse 33, Berlin, W. 30, Germany. (1931.)

296. Tabor, Mr R. J., B.Sc., F.L.S., Botanical Department, Imperial College of Science, South Kensington, London, S.W. 7. (1914.)

297. Telfer, Mr R. Allsop, St Cuthbert's, Malvern. (1931.)

298. Tennessee, University of, Agricultural Experiment Station, Library, Knoxville, Tennessee, U.S.A. (1926.)

299. Tervet, Mr I. W., B.Sc., Edinburgh and East of Scotland College of Agriculture, 13 George Square, Edinburgh.

300. Tetley, Miss U., Quarry Garth, Windermere, Westmorland.

Tomkins, Mr R. G., M.A., Ph.D., Trinity College, Cambridge. (1925.)

302. Topping, Mrs M. P., 3, Southdown Crescent, Cheadle Hulme, Cheshire. (1930.)

303. Tunstall, Mr A. C., Tocklai Experimental Station, Cinnamara P.O., Assam, British India. (1933.)

Vaheeduddin Syed, Department of Plant Pathology, University Farm, St Paul, Minnesota, U.S.A. (1934.)

305. Vanterpool, Mr T. C., M.Sc., Botanical Department, University of Saskatchewan, Saskatoon, Canada. (1929.)

306. Vasudeva, Mr R. S., Cotton Pathologist, Agricultural College, Lyallpur, Punjab, India. (1929.)

307. Venkatarayan, Mr S. V., Senior Assistant Mycologist, Agricultural Department, Bangalore, S. India. (1935.)

308. Wadham, Professor S. M., M.A., Department of Agriculture, The University, Melbourne, Victoria, Australia. (1922.)

309. Wakefield, Miss E. M., M.A., F.L.S., Herbarium, Royal Botanic Gardens, Kew. (1911.)

310. Waldie, Mr J. S. L., B.Sc., C.D.A., Department of Agricultural Botany, The University, Reading. (1925.)

311. Wales, University College of, Librarian, Botanical Department, Aberystwyth, North Wales. (1927.)

312. Wallace, Mr E. R., Agricultural Institute, Kirton, nr Boston, Lincs. (1934.)

313. Wallace, Mr G. B., B.Sc. (Agric.), Ph.D., Department of Agriculture, Morogoro, Tanganyika Territory, East Africa. (1928.)

314. Wallace, Mrs G. B., B.Sc., Morogoro, Tanganyika Territory,

East Africa. (1924.)

315. Wallis, Mr A., Westacre, Station Road, Kettering. (1921.)

316. Ware, Mr W. M., D.Sc., South-Eastern Agricultural College, Wye, Kent. (1924.)

317. Washington, Library, State College of, Pullman, Washington, U.S.A. (1924.)

318. Waterston, Mr J. M., B.Sc., 113, Marchmont Road, Edinburgh. (1934.)

319. Watson, MrW., D.Sc., A.L.S., Cedene, Cheddon Road, Taunton. (1933.)

320. Westerdijk, Professor Johanna, Javalaan 4, Baarn, Holland. (1923.)

321. Western, Mr J. H., B.Sc., Department of Agricultural Botany, University College of Wales, Aberystwyth. (1934.)

322. Weston, Mr W. A. R. Dillon, M.A., School of Agriculture, Cambridge. (1923.)

323. Whetzel, Professor H. H., M.A., New York State College of Agriculture, Cornell University, Ithaca, N.Y., U.S.A. (1914.)

324. Whitaker, Mr F. Owen, 51, Grosvenor Avenue, Carshalton, Surrey. (1921.)

325. Whitehead, Mr T., D.Sc., A.R.C.S., University College of North Wales, Bangor. (1920.)

326. Wilkins, Mr W. H., M.A., D.Ph. Department of Botany, The University, Oxford. (1928.)

327. Williams, Mr P. H., 4, Belmont Villas, Windmill Lane, Cheshunt, Herts. (1930.)

328. Wilson, Miss A. P., M.B.E., A.R.C.S., 116, Fellows Road, London, N.W. 3. (1929.)

329. Wilson, Mr Alastair R., B.Sc., The Botany School, Cambridge. (1933.)

330. Wilson, Mr Malcolm, D.Sc., A.R.C.S., F.L.S., Royal Botanic Garden, Edinburgh. (1921.)

331. Wiltshire, Mr S. P., M.A., Imperial Mycological Institute, Ferry Lane, Kew, Surrey. (1920.)

- 332. Wisconsin, The Library, University of, Madison, Wis., U.S.A. (1923.)
- 333. Wolf, Mr B. L., N.D.A., Cornwall Buildings, 45, Newhall Street, Birmingham. (1923.)
- 334. Wood, Mr F. C., 20, South Farm Road, Worthing, Sussex. (1935.)
- 335. Woodcock, Mr A. J. A., M.Sc., F.E.S., Rhianva, 65, Rock Avenue, Gillingham, Kent. (1926.)
- 336. Woodward, Mr R. C., Ph.D., Imperial Chemical Industries, Ltd., Millbank, London, S.W. 1. (1924.)
- 337. Woolhope, The Naturalists' Field Club, Hereford. (1896.)
- 338. Worcestershire Naturalists' Field Club, c/o Mr W. J. Else, Victoria Institute, Worcester. (1921.)
- 339. Wormald, Mr H., D.Sc., A.R.C.S., Research Station, East Malling, Kent. (1921.)
- 340. Yale University, Library, New Haven, Connecticut, U.S.A. (1930.)
- 341. Yeoman, Mr J. B., M.D., Norton, Wirral, Cheshire.
- 342. Young, Miss Elaine M., Ph.D., M.Sc., c/o The Forest Department, Knysna, Cape Province, South Africa. (1927.)
- 343. Zundel, Dr G. L. I., Botany Building, Pennsylvania State College, State College, Pa., U.S.A. (1929.)
- 344. Zürich, Switzerland, Botanical Garden and Museum, c/o Dr A. U. Däniker. (1921.)
- 345. Zürich, Institut für Spezielle Botanik der Eidg. Techn. Hochschule. (1928.)



RULES

Society's Name and Objects

1. The Society shall be called "The British Mycological Society", and its object shall be the study of Mycology in all its branches.

Members of Society

2. The Society shall consist of Honorary Members, Foundation Members, and Ordinary Members; the number of Honorary Members shall be limited to 20, and that of Foundation Members to 100, but the number of Ordinary Members shall be unlimited.

Honorary Members

3. Honorary Members shall be persons of pre-eminence in Mycology, or who have rendered special service to the Society.

Foundation Members

4. Foundation Members shall be those Members or Societies who joined the Society previous to the limit of 100 Members having been attained.*

Officers

5. The Officers of the Society shall consist of a President, one or more Vice-Presidents, Treasurer, Secretaries and Editor or Editors. They shall be elected annually, at the Annual General Meeting of the Society.

Government of Society

6. The government of the Society shall be vested in a Council, which shall consist of the President and other Officers for the time being, together with two or more other Members who shall be elected annually at the General Meeting, and one-half of whom shall retire each year and not be eligible for immediate re-election. The Members to retire shall be those who have been longest in office or, in case of equality, shall be determined by ballot. Ex-Presidents are ex-officio Members of the Council.

Every Meeting of the Council shall be duly summoned by the Hon. Secretary by at least seven days' notice in writing to each Member of the Council.

Period of Office

- 7. The Officers and Council shall hold office as from the 1st of January following their election.
 - * The limit of 100 Foundation Members was reached 22nd October, 1903.

Plant Pathology Committee

8. The special interests of Plant Pathology shall be delegated to an executive committee, to be called the Plant Pathology Committee of the British Mycological Society. This Committee shall consist of the President and Secretaries ex-officio and twelve other members of the Society. The latter shall be elected annually at the Annual General Meeting of the Society, and one-quarter shall retire in rotation each year and shall not be eligible for immediate re-election. The members to retire shall be those who have been longest in office, or, in case of equality, shall be determined by ballot.

The Officers shall consist of a Chairman and a Secretary, to be

elected by the Committee each year.

At least two meetings shall be held every year, six members to

form a quorum.

The Committee shall have power to appoint for any special purpose a sub-committee consisting either wholly or partly of members of the Committee.

Election of Members

9. Honorary Members shall only be elected at a Meeting of the

Society by a majority of the Members then present.

All Ordinary Members shall be proposed and seconded respectively by existing Members, who shall sign a certificate (see Appendix) of recommendation, one at least of the proposers so certifying from personal knowledge. Every candidate for election shall sign an undertaking to abide by the Rules if elected (see Appendix). They shall be elected by a majority of the Members present at any meeting of the Society or by the President and Officers of the Society.

Subscription

10. All Ordinary Members and Societies shall pay an annual subscription of one pound, and Foundation Members five shillings, due on the 1st of January in each year. Honorary Members shall be exempt from any annual subscription.

Any Member wishing to retire from the Society shall give notice to the Hon. Secretary or Treasurer in writing before the 1st of

December of the previous year.

Meetings

11. The Society shall hold one or more meetings annually, at a place and time determined by the Members at the preceding Annual General Meeting, or by the Council. The Annual General Meeting, for the election of officers and the transaction of other business, shall coincide with the Autumn Foray.

Accounts

12. At the Annual General Meeting of the Society in each year the Hon. Treasurer shall present duly audited accounts.

Alteration of Rules

13. The Rules shall not be altered except by a two-thirds majority of the Members present at an Annual General Meeting. A printed copy shall be sent to every Member of the Society on election, and in the event of alterations to all Members.

APPENDIX

Form of proposal for Ordinary Membership of the British Mycological Society

of		
Mycological Society certify that we con	y, we, the undersigne	ry Member of the Britis d Members of the Society desirable Member of the r election.
Dated this	day of	19
	(From per	sonal knowledge.)
-		

Certificate to be signed by the Candidate

I hereby certify that I desire to become an Ordinary Member of the British Mycological Society and that I will abide by the Rules if elected.



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